## PHASE RELATIONS AND MAGNETIC PROPERTIES OF MM-Co-T (T=Fe and Cu) ALLOYS BETWEEN 2:7 AND 2:17 STOICHIOMETRIES

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INTERDISCIPLINARY PROGRAMME IN MATERIALS SCIENCE
INDIAN INSTITUTE OF TECHNOLOGY KANPUR
AUGUST, 1981

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### PHASE RELATIONS AND MAGNETIC PROPERTIES OF MM-Co-T (T=Fe and Cu) ALLOYS BETWEEN 2:7 AND 2:17 STOICHIOMETRIES

A Thesis Submitted
in Partial Fulfilment of the Requirements
for the Degree of

DOCTOR OF PHILOSOPHY

By
EMUR MANICKAM THANGAVELU

to the

INTERDISCIPLINARY PROGRAMME IN MATERIALS SCIENCE
INDIAN INSTITUTE OF TECHNOLOGY KANPUR
AUGUST, 1981

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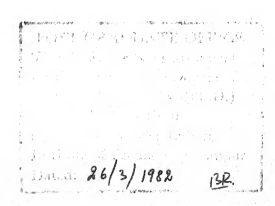
#### CERTIFICATE

Certified that this work on "PHASE RELATIONS AND MAGNETIC PROPERTIES OF MM-Co-T (T = Fe and Cu) ALLOYS BETWEEN 2:7 AND 2:17 STOICHIOMETRIES" by E.M.T. Velu has been carried out under my supervision and that this has not been submitted elsewhere for a degree.

Tehly

August, 1981.

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#### LIST OF SYMBOLS

A, KA : Ampere, Kilo Ampere

a/o : Atom percent

B : Magnetic induction

Br : Remanent induction

(BH) max : Maximum energy product

cc : Cubic centimeters

cps : Counts per second

emu : Electromagnetic unit

g, gms : gram, grams

G, KG : Gauss, Kilo Gauss

H : Magnetic field

H<sub>A</sub> : Magnetocrystalline anisotropy field

HC, bHC: Intrinsic coercivity, induction coercivity

J : Joule

k<sub>1</sub> : Magnetocrystalline anisotropy constant

m : Meter

MM : Mischmetal

MGOe : Million Gauss-Oersted

 $M_{o}$ ,  $4\pi M_{o}$ : Saturation magnetisation

Oe, KOe : Oersted, Kilo Oersted

R. RE : Rare earth

t : Time

T : Tesla (104 Gauss)

w/o : Weight percent

μ : Micron

Investigations were undertaken (i) to establish the relation between structure, microstructure and composition of mischmetal (Indian)-cobalt-iron and mischmetal-cobalt-copper systems in the technologically important composition region and (ii) to develop the technology necessary to produce permanent magnets utilising the commercial mischmetal. The work is divided into eight chapters.

Chapter 1 gives a brief review of the existing literature on the phase diagrams and phase relationships of the binary RE-Co, RE-Fe, RE-Cu and the ternary Sm-Co-Cu alloy systems and an up-to-date critical survey of the developmental processes and properties reported for Rare-Earth Cobalt permanent magnets containing mischmetal.

Chapter 2 is the statement of the problem. The objectives of the work were (i) to investigate the phase relationships in the MM-Co-Fe and MM-Co-Cu alloy systems in the composition region of interest for permanent magnets (ii) to characterise the primary magnetic properties of the MM-Co-Fe and MM-Co-Cu phases in order to evaluate their suitability for permanent magnets and (iii) to study and optimise different process parameters to produce MM-Co permanent magnets.

Chapter 3 describes the experimental procedures followed in the phase analysis and the magnet fabrication.

Alloys for the study were prepared by arc melting in a water cooled Cu crucible under Argon atmosphere and subsequently homogenised at 900°C in evacuated quartz ampoules. The phases were studied by metallography, X-ray diffraction, thermomagnetic analysis and electron-probe micro-analysis. Magnets were prepared by powder metallurgy techniques. The as-cast alloy buttons were crushed to-too mesh and ground to a few micron size powder by rod milling under sodium-gettered toluene. The powder was dried under a vacuum and field pressed in 13 KOe pulse field. The powder compacted were sintered under flowing argon atmosphere. The sintered pellets were magnetised in a pulse field of 60 KOe and their permanent magnet properties were evaluated with a vibrating sample magnetometer.

In Chapter 4 the results of phase relations study of the MM-Co-Fe system at 900°C are discussed. In MM-Co system, six phases, MMCo<sub>3</sub>, MM<sub>2</sub>Co<sub>7</sub>, MM<sub>5</sub>Co<sub>19</sub>, MMCo<sub>5</sub>, MM<sub>2</sub>Co<sub>17</sub> and β-Co were identified between 72 and 92 a/o Co. The MMCo<sub>3</sub>, MM<sub>2</sub>Co<sub>7</sub> and MM<sub>5</sub>Co<sub>19</sub> have very similar X-ray patterns and hence they were identified by thermomagnetic analysis (TMA) based on their distinct Curie temperature. The MM<sub>2</sub>Co<sub>7</sub> phase exists both in the hexagonal and rhombohedral modifications. The 2:7, 5:19, 1:5 and 2:17 phases also exist as stable phases at 900°C when Co is partially substituted by Fe. Alloys containing upto 20 a/o Fe were investigated in close composition interval of 1 a/o. The 5:19 phase was found to

extend up to about 11 a/o Fe while the 2:7 and 1:5 phases extend to greater than 11 a/o Fe. A new phase with a T of 340°C was found to coexist with a number of alloys near the 1:4 composition. A 900°C isothermal section of the ternary MM-Co-Fe system between 2:7 and 2:17 stoichiometry (MM/Co) is given with alloys containing upto 20 a/o Fe. The phase region beyond 12 a/o Fe in the MM-Co-Fe system particularly between 2:7 and 1:5 stoichiometric regions seems to be complicated because the X-ray results alone could not establish the identity of the phases unambiguously. Some of the phases in this region have T greater than the safe-limit (700°C) of the TMA apparatus used in this study. Of the various phases only MMCo5 has an observable homogeneity region extending over 2 w/o Co at 900°C. The MMCo5 phase is unstable at temperature below 700°C and the addition of Fe increases its instability.

In Chapter 5 the primary magnetic properties, saturation magnetisation and Curie temperature of the various intermetallic phases identified in the MM-Co-Fe system are reported. The 1:3, 2:7, 5:19, 1:5 and 2:17 phases in the MM-Co system have  $T_{\rm C}$  <20°C, 67°C, 280°C, 540°C and >700°C respectively. The substitution of Fe for Co increases these values further for all these phases. The MMCo<sub>5</sub> has a  $4\pi$  M<sub>S</sub> of 95 emu/g at 20°C and that of MM<sub>2</sub>Co<sub>17</sub> phase is 114 emu/g. Both the 5:19 and 2:7 phases have a  $4\pi$  M<sub>S</sub> less than 45 emu/g.

Chapter 6 contains exclusively results on phase relations and magnetic properties of MM( ${\rm Co}_{1-x}{\rm Cu}_x$ )<sub>5</sub> 0  $\leqslant$  X  $\leqslant$  1, alloys. Cu can be substituted for Co upto 100% giving rise to stable  ${\rm CaCu}_5$  type phase. At low Cu concentration, 2:17 and a new phase coexist with the matrix 1:5 phase at 900°C. Both the <u>a</u> and <u>c</u> of the hexagonal 1:5 lattice increase with increasing Cu concentration. Cu can be substituted for Co upto 15 w/o in the 1:5 phase with a T<sub>C</sub> not less than 500°C and a  $4\pi$  M<sub>S</sub> of about 70 emu/g. Beyond 15 w/o both the  $4\pi$  M<sub>S</sub> and the T<sub>C</sub> are considerably reduced.

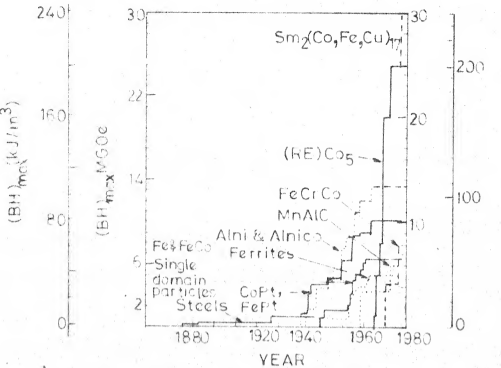
Chapter 7 deals with the permanent magnet properties of MM-Co alloys. Out of the many process parameters for magnet production a selected number of the parameters like, alloy composition, milling time, sintering temperature and time were studied for MMCo<sub>X</sub>, 3.5 £ X £ 4.8 alloys. Powder compacts were sintered between 960 and 1030°C for 5 m to 60 m. A maximum iH<sub>C</sub> of 5250 oe was obtained for MMCo<sub>4.0</sub> alloy sintered at 1000°C for 15 m. A 20% substitution of Sm to MMCo<sub>3.8</sub> alloy increased the iH<sub>C</sub> to greater than 11 Koe. The sintered pellets contained invariably considerable amount of MM<sub>2</sub>O<sub>3</sub> as revealed by X-ray and metallography. The chemical analysis of the powder compact and its sintered pellet showed same composition but it was different from the composition of the as-cast alloy indicating that the oxidation should have occurred during the process of powder production.

Chapter 8 is on the conclusions and suggestions for future work. A summary of the results obtained under Chapters 4, 5, 6 and 7 is given. The problem to be solved in the high Fe containing MM-Co-Fe alloys and the steps to be taken to minimise oxidation and improve the permanent magnet properties of the MM-Co alloys are given.

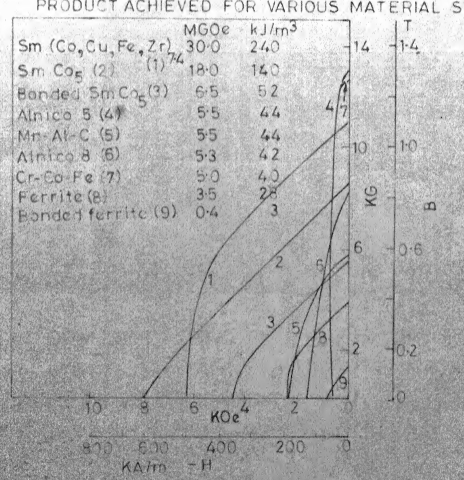
#### 1. INTRODUCTION

#### 1.1 A Comparison of Different Permanent Magnets

A permanent magnet is characterised by its coercivity (H<sub>C</sub>), remanent induction (B<sub>r</sub>) and maximum energy product  $((BH)_{max})$ . High values for H and B give a magnet with a high (BH) max. The (BH) represents the maximum energy storage per unit volume and is the figure of merit often used for permanent magnet materials. In Figure 1.1 the progress in permanent magnet quality since 1880 as indicated by the value of the maximum energy product achieved for various material systems is illustrated 1. It could be seen from this figure that the actual development of RE-Co magnet started around the year 1965 and the progress made within a decade following the year 1965 is spectacular over the erstwhile dominating magnets such as alnicos and ferrites. Rare earth cobalt magnets basically of two compositions, RECos and Sm2(Co,Fe,Cu)17 have been developed. The demagnetisation curves of selected permanent magnet materials are given in Figure 1.2 where it could be seen again that RECos and RE(Co, Cu, Fe, Zr) 17-x alloys have superior demagnetisation curves to those of other magnets being used currently 2. The repelling forces of two identical magnets of alnico, ferrite and a rare earth cobalt magnet (recoma, 1:5 type) is illustrated in Figure 1.3 for a comparison of their strength. 3 The supporting strength of the three materials are in the ratios of 1:6.5:37 respectively.



AS INDICATED BY THE VALUE OF THE MAXIMUM ENERGY PRODUCT ACHIEVED FOR VARIOUS MATERIAL SYSTEMS (Ref. 1).



G.12 DEMAGNETISATION CURVES OF SELECTED PERMANENT MAGNET MATERIALS (Ref. 2)

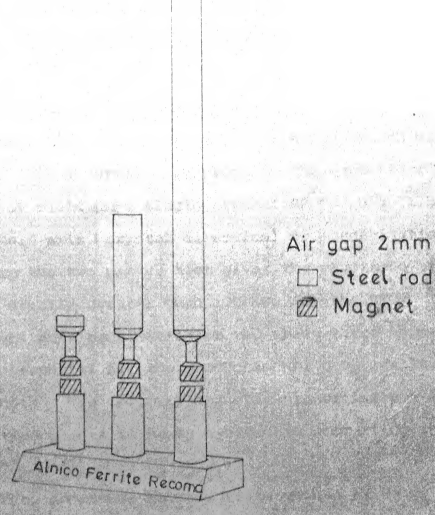


FIG.1.3 DEMONSTRATING THE REPELLING FORCES OF TWO IDENTICAL MAGNETS OF ALNICO 8, STRONTIUM FERRITE AND RECOMA (Ref. 3)

The supporting strength of the three materials are in ratios of 1:6.5:37 respectively.

A Recoma magnet supports 74 times its own weight

#### 1.2 Primary Magnetic Properties of RE-Co Phases

The three primary magnetic properties that determine the suitability of a RE-Co phase for permanent magnet potential are (1) the magnetocrystalline anisotropy constant (K1) (2) saturation magnetisation ( $4\pi M_c$ ) and (3) Curie temperature ( $T_c$ ). The magnetocrystalline anisotropy constant ( $K_1$ ) is related to magnetocrystalline anisotropy field  $(H_{\lambda})$  by  $H_{\lambda}$  =  $2K_1/M_s$  where  $H_A$  is the field necessary to rotate the magnetisation in a perfect single domain crystal. The  ${\rm H}_{\rm A}$  and  ${\rm M}_{\rm S}$ are the experimentally determined quantities from which  $K_1$  is deduced. The  $\mathbf{H}_{\mathbf{A}}$  is determined by plotting the magnetisation as a function of field for a single crystal RE-Co phase along its easy and hard axis (crystal direction) of magnetisation. The point where the two curves meet gives the magnitude of  ${\rm H_A}$  and it is usually greater than 100 KOe for the RECo $_5$  type phases. A favourable combination of the above three magnetic properties is essential for the production of good quality permanent magnets. Then primary magnetic properties for the 1:5 and 2:17 type phases in RE-Co systems are summarised in Table 1.1.

The RECo<sub>5</sub> compounds have moderate values of  $4\pi$  M<sub>s</sub> ( $\sim$ 1 T) but extremely large values of K<sub>1</sub> ( $\gg$ 1 MT/m<sup>3</sup>). The latter property is primarily responsible for the exceptionally large coercivity. In this regard SmCo<sub>5</sub> is the most outstanding of all RECo<sub>5</sub> magnets by virtue of having the largest value of K<sub>1</sub> (17 MJ/m<sup>3</sup>). The RE<sub>2</sub>Co<sub>17</sub> compounds have higher  $4\pi$  M<sub>s</sub>

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TABLE 1.1 MAGNETIC PROPERTIES OF RCo5 AND R2Co17 COMPOUNDS:  $4\pi$  Ms AND K1 VALUES AT 25°C ARE GIVEN

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Rare	t t t	RCo <sub>5</sub>		R <sub>2</sub> Co <sub>17</sub>			
earth	4π M <sub>s</sub>	T <sub>C</sub>	К <sub>1</sub>	4π'M s	T <sub>C</sub>	K <sub>1</sub>	
	(T)	(°C)	(MJ/m <sup>3</sup> )	(T)	(°C)	(MJ/m <sup>3</sup> )	
Ce	0.85	374	5.3	1.15	800	-0.6	
Pr	1.12	612	8.1	1.38	890	-0.6	
Nd	1.20	630	0.7	1.39	900	-1.1	
Sm	0.97	724	17.2	1.20	9 20	3.3	
Gđ	0.19	735	4.6	0.73	930	<b>-0.5</b>	
Tb	0.24	707		0.68	920	-3.3	
Dy	0.30	6930		0.70	910	-2.6	
Но	0.53	727	3.6	0.83	9 20	-1.0	
Er	0.63	713	3.8	0.90	930	0.41	
Tm	0.67	747		1.13	920	0,50	
Yb						-0.38	
Lu				1.27	940	-0.20	
La	0.91	567	5.9				
Y	1.06	648	5.2	1.25	940	-0.34	
Th			2.6			-0,53	

T = 10 KG.

values than the corresponding RCo $_5$  series and hence might have greater energy products. However, as Table 4.1 shows, the magnitude of  $K_1$  for the 2:17 compounds is considerably smaller than that for the corresponding 1:5 series. Furthermore, with the exception of Sm, Er and Tm, all 2:17 binary compounds have  $K_1$  0, that is, an easy (0001) plane. For these compounds, magnetisation reversal becomes relatively easy and low coercivity is expected. There has been some success at modifying the sign and magnitude of  $K_1$  by partial substitution for Co by other transition metals such as Fe, Cr and Mn. $^4$ 

The magnetism of the RE-Co compounds is due to the interatomic exchange between the spins of the two sublattices plus the spin-orbit coupling within the rare earth atoms. In the lighter rare earth series-Ce, Pr, Nd and Sm - the spins of the RE and Co atoms are aligned parallel. The value of  $4\pi$  Mg are thus high. In the other they are aligned antiparallel, and the values of  $4\pi$  Mg tend to be low. Yttrium is nonmagnetic and hence the magnetic induction comes from Co alone. The magnetocrystalline anisotropy also comes from two sources, one originating in the itinerant electrons of the Co sublattice and one due to the crystalline electric field of the rare earths. A broad summary ranging from basic magnetism to the technology of the rare earth magnets is given by Menth et.al.  $^5$ 

## 1.3 RE- 3d Metal Phase Diagrams and Structures of Their Intermetallic Compounds

#### 1.3.1 RE- 3d Metal Phase Diagrams

The binary phase diagrams of Ce-Co, 6,7 La-Co, 8

Ce-Fe<sup>9</sup> and Ce-Cu<sup>7</sup> are shown in Figures 1.4a to 1.4d and the compound formation in the high transition metal side in these systems is shown in Figure 1.5. The phase diagrams of Nd-Co and Pr-Co systems are similar to that of Ce-Co systems. 8 It is seen from Figure 1.4a that all the intermetallic phases in the Ce-Co system form through peritectic reaction. The temperatures of peritectic reaction for the intermetallic phases in the RE-Co systems (RE = Cr, La, Nd and Pr) are shown in Table 1.2.

TABLE 1.2 PERITECTIC TEMPERATURES OF INTERMETALLIC PHASES IN RE-Co SYSTEMS<sup>7,8</sup>

System	Peritectic temperature (°C)				
	1:3	2:7	5:19	1:5	2:17
Ce-Co	1091	1130	1134	1193	1209
La-Co	Does not exist	843	868	1124	Does not exist
Nd-Co	1105	1161	1166	1266	1301
Pr-Co	1057	1118	1125	1229	1271

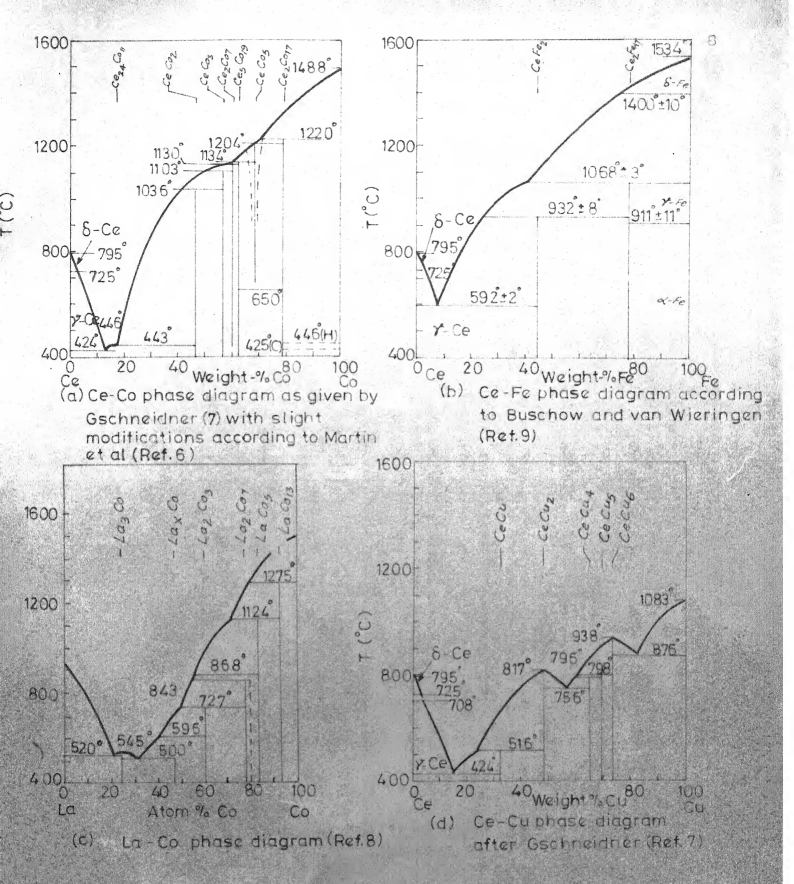
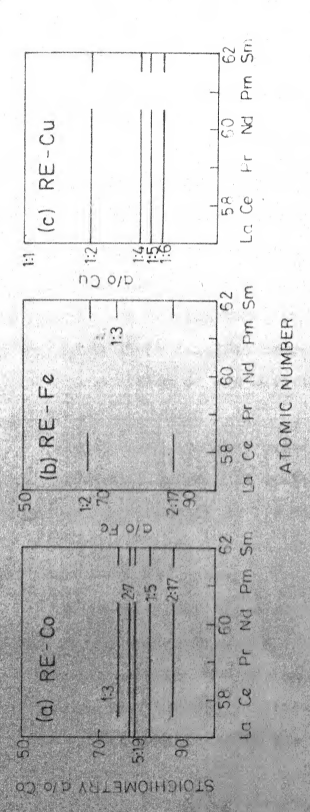


FIG. 1.4 PHASE DIAGRAMS OF RE-T SYSTEMS RE-Ce and La, T-Co, Fe and Cu



COMPOUND FORMATION IN (a) RE-Co (b) RE-Fe AND (c) RE- Cu SYSTEMS FIG. 1.5

In the case of La-Co system the LaCo $_3$  and La $_2$ Co $_{17}$  phases do not exist. Also for the La $_2$ Co $_7$  and La $_5$ Co $_{19}$  phases the peritectic temperatures are lower by about 300°C than that corresponding to these phases in the other RE-Co systems (Table 1.2). In Ce-Co system (Figure 1.4a) two features could be noted for the 1:5 phase. It shows a homogeneity region at high temperatures ( $_3$ 700°C) and undergoes an eutectoid type reaction at temperature below about 650°C. This was observed for other RE-Co systems also and the eutectoid decomposition temperatures for various RECo $_5$  phases are shown in Figure 1.6 together with their liquidus temperatures. The effect of Fe and Al impurities on the eutectoid reaction temperatures is also shown in Figure 1.6.

In the case of Ce-Fe system only 1:2 and 2:17 intermetallic compounds exist. In Sm-Fe system in addition to these phases, the existence of an 1:3 phase is reported. The 2:7, 5:19 and 1:5 phases found in the RE-Co systems are absent in the RE-Fe system.

In Ce-Cu system<sup>7</sup> five intermetallic phases corresponding to the stoichiometries, 1:1, 1:2, 1:4, 1:5 and 1:6 are seen, out of which the 1:2 and 1:6 phases form congruently and the rest form through peritectic reaction. The 2:7, 5:19 and 2:17 phases found in the Ce-Co system do not exist in this system.

Besides the binary systems, the isothermal sections of Sm-Co-Cu phase diagram reported by Perry et.al. 13 at 800°C

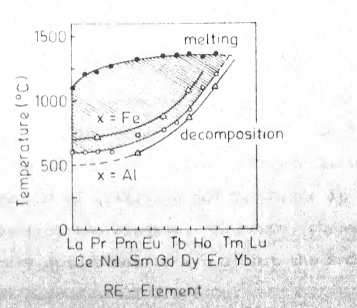
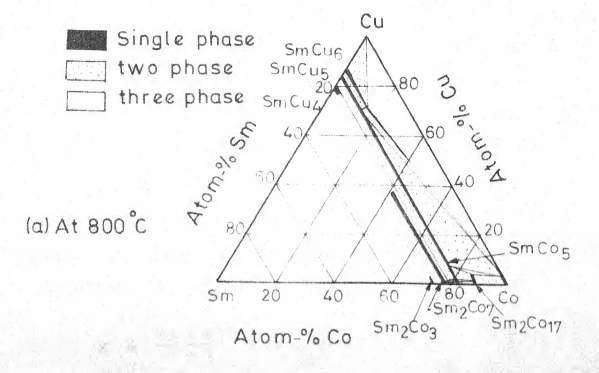


Fig 1.6 Melting temperature and decomposition temperature for various members of the RECog compound. The curves denoted by X = Fe and X = Al show the decrease or increase of the % stable CaCus - type region that can be obtained by adding 3 at % X as an impurity (Ref 11)

1200°C are shown in Figures 1.7a and 1.7b. It is seen from Figure 1.7a that at 800°C, the 1:5 phase could exist as a stable phase with 100% solid solubility for Cu. The Sm<sub>2</sub>Co<sub>7</sub> phase extends into the ternary upto about 38 a/o Cu but the SmCo<sub>3</sub> and Sm<sub>2</sub>Co<sub>17</sub> phases extend only upto a few a/o Cu into the ternary. Two narrow three-phase regions, one containing the 1:5, 2:17 and Co phases and the second containing 2:7, 1:5 and 2:17 phases are present close to the Sm-Co binary line. These regions do not exist at 1200°C (Figure 1.7b). A third three-phase region of 2:17, 1:6 and Co phases is present in the Cu rich part of the diagram. The SmCu<sub>6</sub> phase extends into the ternary upto about 16 a/o Co but the SmCu<sub>4</sub> phase does not exist beyond about 2 a/o Co.

#### 1.3.2 Structures of Intermetallic Phases in RE-Co Systems

from the REM<sub>2</sub> to the RE<sub>2</sub>M<sub>17</sub> intermetallics (RE = Ce, Nd, Pr and Sm, M = Co). The compounds in this range can be arranged into two groups, hexagonal and rhombohedral, which derive from the hexagonal CaCu<sub>5</sub> type structure of the REM<sub>5</sub> alloys. 14-17 The structure of the 1:5 alloy consistsof an infinite stacking sequence of two-atom layers in the direction of c-axis: a hexagonal arrangement of M atoms followed by a mixed layer of RE and M atoms (Figure 1.8a). 18 The hexagonal unit cell contains a three rhombohedral unit cells (Figure 1.8b). 14



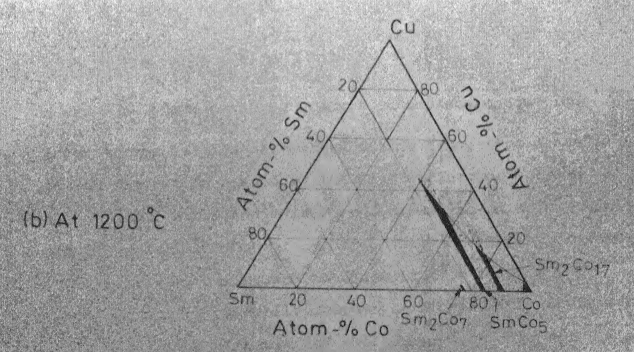


FIG. 17 ISOTHERMAL SECTIONS OF THE Sm-Co-Cu PHASE DIAGRAM AFTER PERRY(Ref. 13)

first, second, third etc. unit cell, a simultaneous shift of the newly formed building blocks in the hexagonal basis and minor rearrangements of the RE atoms, the compounds  $\text{REM}_2$ ,  $\text{REM}_3$ ,  $\text{RE}_2^{\text{M}}_7$  etc. are formed.  $^{14,15,17}$  The respective atom positions for  $\text{REM}_3$  are shown in Figure 1.8c; the mode of formation of the other compounds is schematically given in Figure 1.9. The stoichiometries of the possible intermetallics in the series  $\text{REM}_{\chi}$  can be calculated from

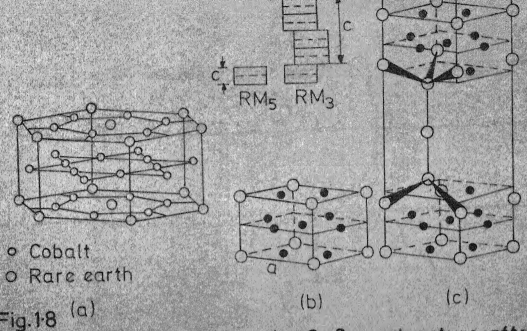
$$X = \frac{5n+4}{n+2}$$
  $(n = 0, 1, 2, 3 ....)^{15}$ 

and the theoretically possible Cromer-Larson phases in RE-Co system are shown in Table 1.3. The lattice parameters of these phases are given by 19,20

$$a_0 = a_0 (CaCu_5) + d_a (n + 1)$$

$$c_0 = m(n + 1) c_0 (CaCu_5) + d_c/(n + 1)$$

where  $a_0$  (CaCu<sub>5</sub>) and  $a_5$  (CaCu<sub>5</sub>) in the above equations. refer to the lattice parameters of the CaCu<sub>5</sub> type phase, d is the distortion caused by the size difference between RE and 3d metal atoms and m is 2 or 3 depending on wether the structure is rhombohedral or hexagonal. From  $RE_2M_7$  onwards (Table 1.3) the difference in concentration between the predicted phases gradually decreases; finally this difference becomes indistinguishable and the series converges for  $n = a_2$  against the  $RECo_5$ . Strnat et.al. Thave reported thermomagnetic



- a) Hexagonal unit cell of the CaCu<sub>5</sub> structure after (Ref. 18)
- b) Rhombohedral unit cell of the CaCu<sub>5</sub> structure after (Ref. 14) (c) Substitution of one RE against one M atom in every second
- CaCus unit cell minor displacements of RE atoms and layer shifts in the hexagonal basis plane lead to the REM3 structure (Ref. 14)

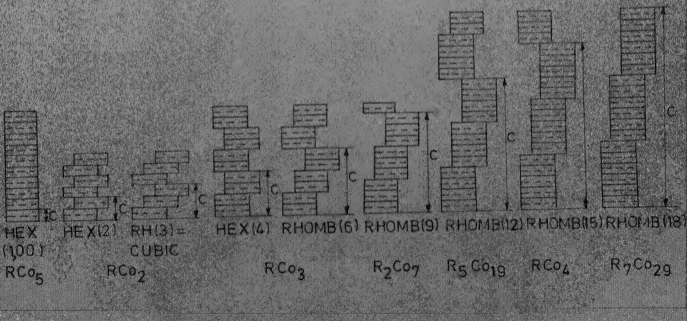


Fig. 1.9 Schematic diagram of the mode of formation of hexagonal and rhombohedral R<sub>x</sub> Co<sub>y</sub> alloys out of the RCo<sub>5</sub> structure by layer shifts in the hexagonal base plane after (Ref. 14,15,17)

TABLE 1.3 THEORETICALLY POSSIBLE CROMER-LARSON PHASES IN RE-Co SYSTEMS

n	Phase RE <sub>x</sub> Co <sub>y</sub>	Co/RE ratio
0	1:2	2
1	1:3	3
2	2:7	3.5
3	5:19	3.8
4	1:4	4.0
5	7:29	4.14
6	4:17	4.25
7	3:13	4.33
• • •	• • •	• • • •
104	1:4.99	4.99
• • •		• • •
00	1:5	5.0

evidence for the phases of stoichiometry 1:4, 7:29, 4:17 and 3:19 in Pr-Co system but no X-ray or microstructural evidence is reported for these phases.

## 1.4 Structure Sensitive Magnetic Properties

The magnetocrystalline anisotropy  $^{21}$  field  $_{
m A}$ , and the coercivity  $^{22}$  of the RECo $_5$  alloys depend on the microstructure of the alloy. It is seen from Figure 1.10 that the  $_{
m A}$  of 1:5 phase peaks at composition corresponding to 1:5 stoichiometry and falls steeply on either side where the 1:5 coexists with either 2:7 or 2:17 phases. In Figure 1.11 the dependence of  $_{
m i}^{
m H}_{
m C}$ ,  $_{
m b}^{
m H}_{
m C}$  and (BH) on the composition of Sm-Co alloy is shown. The coercivity peaks on the slightly Sm richer side of SmCo $_5$  phase but it drops steeply on the Co rich side of 1:5 composition. This indicates that the presence of small amount of 2:7 phase with the 1:5 phase is not detrimental to  $_{
m i}^{
m H}_{
m C}$  as the presence of 2:17 phase with the 1:5 phase.

In Figure 1.12 the effect of heat treatment on the  ${}_{^{1}\text{C}}$  of  $\text{SmCo}_{5}$  magnet is shown. When the  $\text{SmCo}_{5}$  magnet is heated around  $750^{\circ}\text{C}$  for 30 minutes, the  ${}_{^{1}\text{C}}$  falls from about 35 KOe to about 5 KOe. This drastic drop in  ${}_{^{1}\text{C}}$  is attributed to an eutectoid reaction in  $\text{SmCo}_{5}^{23}$  where the 1:5 phase decomposes into the adjacent 2:7 and 2:17 phases.

- 1.5 Technology of RE-Co Magnet Processing and Permanent Magnet Properties of RECo<sub>5</sub> Type Magnets
- 1.5.1 Flow Chart of RE-Co Magnet Processing

Powder metallurgy techniques are employed in the manufacture of rare-earth cobalt permanent magnets. The basic process consists of preparing powder by one of several

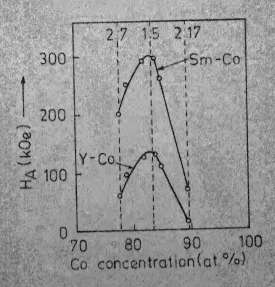


Fig. 1-10 Magnetocrystalline anisotropy of RE - Co materials as a function of composition (Ref 21)

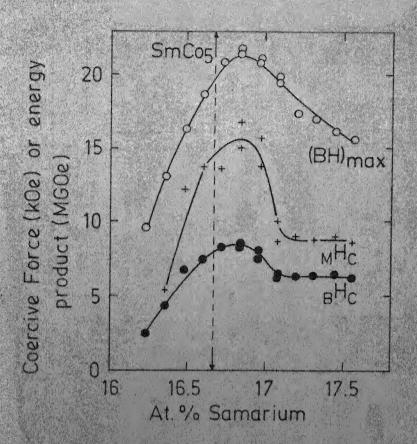


Fig. 1:11 Dependence of intrinsic as well as induction coercivity,  $_{\rm M}H_{\rm C}$  and  $_{\rm B}H_{\rm C}$  and energy product,  $({\rm BH})_{\rm max}$ , of SmCo5  $\pm$  x magnets on cobalt content. (Ref 22)

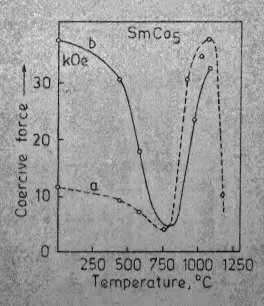


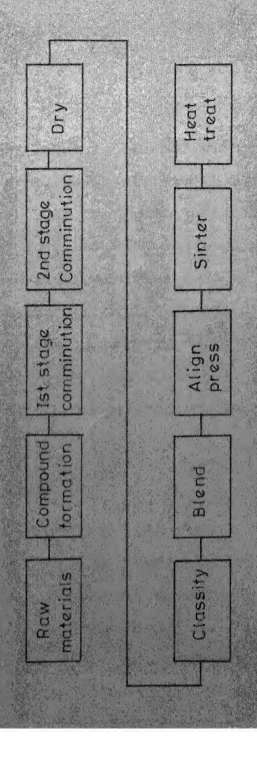
Fig. 1.12 Coercivity measured at room temperature as a function of heating temperature; samples were heated for 30 minutes; reaction with oxygen was avoided. Curve a ground material Curve b ground material heated at 1080°C; second time heated at indicated temperature (Ref. 23)

methods, pressing the powder in a magnetic field, sintering the compacts and heat treating the sintered product. The overall process flow diagram is illustrated in Figure 1.13. Close control is required throughout and the most important factors are chemical composition, oxygen content, powder particle size distribution, aligning field and sintering and heat treating cycles.

## 1.5.2 Magnet Processing Parameters

## 1.5.2.1 Alloy Compositions

The chemical composition of the alloy determines its phase composition and the properties of the magnet produced depends on its microstructure. Therefore, a close control of the chemical composition of the alloy is necessary during magnet processing. The change in composition can occur during melting due to the loss of RE elements by evaporation or during powder preparation and sintering due to oxidation of RE elements. Thus the final composition of the magnet may turn out to be different from the starting composition of the alloy. The extent of shift in composition will depend on the melting method (arc or induction melting), milling procedure, and sintering process (solid or liquid phase sintering). Thus for each material system and for each set of process parameters adopted, the composition shift may be different. Therefore the right composition of the sintered magnets that gives best properties is to be determined by a



PROCESS FLOW DIAGRAM FOR SINTERED RARE EARTH COBALT PERMANENT MAGNETS (Ref. 26) FIG. 1:13

systematic study of these variables. It has always been reported that in the production of  ${
m RECo}_5$  type magnets a RE-rich composition is used.  $^{24,25}$ 

## 1.5.2.2 Comminution

The cast ingot is crushed and ground to produce a powder with the proper size distribution and flow characteristics. In Figure 1.14 the proper particle size distribution (dashed curve) is compared with one that is too bad (solid curve). The large-size particles tend to be multi-grained and do not align or sinter properly. Under sized particles pick up an excessive amount of oxygen, which reacts with the metallic components to form oxide resulting in an overall shift of the composition out of the range where optimum properties are achieved.

## 1.5.2.3 Compaction

The properly sized and dried powder is pressed in the presence of an applied magnetic field not less than 10 KOe. 26 Homogeneous fields are necessary to produce a high level of uniformly aligned particles. It has been found that the application of a field in a direction perpendicular to the pressing direction leads to a better alignment of particles. Finished magnets are anisotropic with the direction of magnetisation determined during the pressing aligning step.

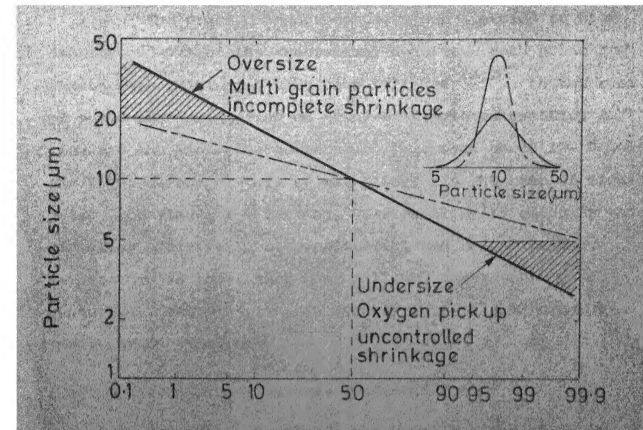


FIG. 1.14 COMPARISON OF OPTIMUM PARTICLE SIZE
DISTRIBUTION (DASHED CURVE)
WITH BROAD PARTICLE SIZE DISTRIBUTION
(SOLID CURVE)(Ref. 26)

## 1.5.2.4 Sintering

The pressed powder compacts are sintered to high density, 93-97% of the theoretical density. Both solid and liquid phase sintering have been followed. 28,29 In the case of solid phase sintering an alloy of single composition is used and the sintering temperature is chosen about 100°C below the liquidus temperature of the alloy. In the case of liquid phase sintering a low melting, RE-rich alloy is added as the sintering additive to a base material and the sintering temperature is lower than that followed for a solid phase sintering. The sintering is usually carried out in a high purity argon atmosphere.

## 1.5.2.5 Post-sintering Heat Treatment

temperature (<900°C) to optimise overall second quadrant characteristics of the hysteresis loop. 29-31 The shape of the demagnetisation curve as well as the magnitude of the coercivity can be considerably altered by suitable heat treatment procedure. 31 This heat treatment is found to be associated with change in microstructure of the magnets.

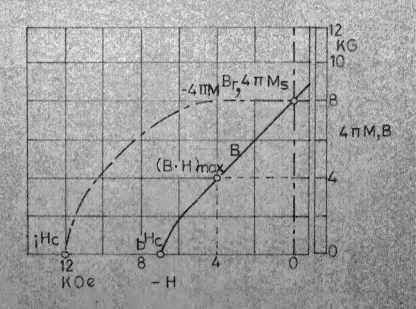
## 1.5.2.6 Magnetising and Characterisation

The permanent magnet properties are characterised by measuring its hysteresis curve particularly its second quadrant (demagnetisation curve). Two methods of representation are used; the intrinsic magnetisation curve and the BH

hysteresis curve (Figure 1.15). The intrinsic magnetisation curve is more suited to describing the physical properties of the material while the BH curve is used for all practical purposes. The latter curve can be obtained from the former by the relation  $B = H + 4\pi$  M. The  $B_r$ ,  $bH_c$  and  $(BH)_{max}$  are obtained directly from the demagnetisation curve. The field applied for magnetising the samples during measurement should be sufficiently high to saturate the material, otherwise there is the risk of tracing the minor hysteresis loop. The field level is normally determined by the intrinsic coercivity in the case of RECo<sub>5</sub> type magnets. In practice the magnets are magnetised prior to magnetic measurements in a pulse field of 60-100 KOe and then their demagnetisation curves are traced in a magnetic field sufficiently higher than the intrinsic coercivity of the magnet.

1.5.3 Processes and Properties of Selected RECo<sub>5</sub> Type Magnets

Procedures followed in magnet processing and the properties developed for a selected number of RECo5-type magnets are summarised in Table 1.4.



## FIG.1-15 TWO REPRESENTATIONS OF THE HYSTERESIS CURVE OF A PERMANENT MAGNET (Ref. 3.)

4 TM = Intrinsic magnetization curve

B = Induction curve

The relationship between the two curves is given by

B = 4 T M + H

in the Gaussian system of units

411 M= Saturation magnetization

Br = Remanence

iHc, bHc = Coercive tield of magnetization and induction respectively

TABLE 1.4 PROCESSES AND PROPERTIES OF SELECTED RECOS TYPE MAGNETS

System	Process	Properties	Reference
SmCo <sub>5</sub>	(a) Induction melted, ground to 25 L size particles by vibrating grinders in toluene. Compacted at 10 KOe. Sintered 1000°C for 60 m in an inert atmosphere.	Br: 9000 G bHc: 9000 G i <sup>H</sup> c: 25000+ 0e (BII) max: 20 MG0e	D. Das <sup>28</sup>
	Sinter additive: 60% Sm, 40% Co Base material: 66.7% Co. 33.3% Sm Induction melted, Fluid energy— milled to 6-8 L. size in nitrogen. Average composition of the biended powder: 62.6% Co. 37.4% Sm. Axially aligned at 60-100 Koe at 200 Kpsi to 6.8 g/cc sintered for 30 m at 1100°C in high purity Argon to 7.7 g/cc	4π M <sub>S</sub> : 9.4 KG B <sub>L</sub> : 8.1 KG bH <sub>C</sub> : 7.7 KOe i H <sub>C</sub> : 16.4 KOe (BH) <sub>max</sub> : 15.7 MGOe	M.G. Benz 29 et.al.
Sm <sub>O,5</sub> MM <sub>O,5</sub> Co <sub>5</sub>	Base material: 66 w/o Co, 17 w/o MM and 17 w/o Sm. Additive: 40 w/o Co, 60 w/o Sm. Induction melted. Fluid energy—milled to ~10 % in nitrogen. Aligned in 60 Koe field and compacted at 200 K psi. Sintered at 1000-1075°C for 30-60 m and magnetised in 100 Koe field prior to measurement	$4\pi M_{\rm s}$ : 9.6 KG $_{\rm r}^{\rm B}$ : 9 KG $_{\rm r}^{\rm B}$ : 7.6 KOe $_{\rm r}^{\rm H}$ : 15.2 KOe (BH) $_{\rm max}$ : 20 MGOe	M.G. Bons et. al 25

TABLE 1.4 (contd.)

System	Process	Properties	Reference
MMCo <sub>5</sub> (MM: 53 a/o Ce, 30 a/o La, 13 a/o Nd and 4 a/o Pr)	Induction melted in Argon atmosphere. Alloys with MM richer than in MMCo5 were jet-milled in nitrogen. Powder aligned in 50 KOe field at 6000 atmosphere. Sintered at 1040°C, heat treated at	i <sup>H</sup> c : 9 KOe B : 8.1 KG (BH) <sub>max</sub> : 14 MGOe	H. Nagel et.al.
MM(Co,Cu,Fe, Mg)5 (MM: 52.1-57.8 w/o Ce, 23.4- 33.4 w/o La, 10.8-17.9 w/o Nd and 3.4- 6.6 w/o Pr)	MM(Co,Cu,Fe, 35.5 w/o MM, 53.5 w/o Co, 6 w/o Fe, 4.9 m/o Cu and 0.1 w/o Mg. Alloys milled in spex shatter box at -60°C in Toluene to spex shatter box at -60°C in Toluene to spex shatter box at -60°C in Toluene to spex size, aligned in a magnetic field w/o Ce, 23.4 of 8 koe and compacted at 45,000 psi. 10.8=17.9 w/o Nd and 3.4 of sintered in He at 980°C for 2 hours. Nd and 3.4 of sintered in He at 980°C for 2 hours.	i c : 5.6 KOe bh : 4.4 KOe Br : 6.9 KG (BH) max : 10.3 MGOe	J.W. Walkiewicz et.al. 32 e

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## 2. STATEMENT OF THE PROBLEM

The review given in Chapter 1 under Section 1.1 shows that permanent magnets of outstanding technical properties  $(_{i}^{H}_{C} \text{ and } (_{BH})_{max})$  much superior to other types of permanent magnets could be produced from Rare-Earth Cobalt alloys. Among the family of Rare-Earth Cobalt magnets, alloys containing Sm as the RE have maximum values for permanent magnet properties. But at the same time Sm is costly and this hinders the large volume production of its magnets for widespread application. In many of the permanent magnet applications a high resistance to demagnetisation is not necessary. In such cases a material which is cheap and at the same time has relatively large values for  $_{i}^{H}_{C}$  and  $_{i}^{G}_{C}$  and  $_{i}^{G}_{C}$  compared to the conventional commercial permanent magnets such as alni $_{i}^{G}_{C}$  and ferrites, may be preferred.

While the technology of production of SmCo<sub>5</sub> magnets has been widely studied and reported, (1-8) only a few studies have been reported on the development of magnets utilising mischmetal (a cerium rich mixture of light Rare-Earths) as the only RE. (9-11) Also the composition of mischmetal differs depending on the mineral source and the extraction method. In India there is a vast deposit of Rare-Earth minerals (in Travancore beach sand, Kerala) and it is the third largest producer of Rare-Earth in the world. The Indian mischmetal employed in this study contains about 6 w/o Fe as impurity introduced during the electrolytic process. Cobalt is less

abundant than RE and is a scarce material. 12 Its partial replacement with other transition elements like Fe and Cu would certainly be preferred if the loss in magnetic properties as a result of their introduction is not much. Hence a study was undertaken to develop permanent magnets utilising the cheap commercial grade Indian mischmetal and cobalt with partial addition of Fe and Cu in place of Co.

The present study was undertaken with two broad objectives. Firstly to study the relation between structure, microstructure and composition and establish the phase relationships in the MM-Co-Fe and MM-Co-Cu systems. Secondly to select the suitable phase compositions in these systems and study the influence of various process parameters on the permanent magnet properties.

Magnetic properties (iH<sub>C</sub> and Br) are very sensitive to phase composition 13-16 and a knowledge of phase relationship in the RE-Co system is essential to select the right composition for the magnet production. Phase relations study in the RE-Co system is complicated because of the close structural similarity among the various intermetallic phases. 17-19 In addition to the conventional experimental tools like metallography and X-ray diffraction, additional experimental techniques such as thermo-magnetic analysis (TMA) and electron probe micro-analysis (EPMA) are imperative to confirm the identity of the phases. Some of the phases like 5:19 in the binary RE-Co system were discovered only lately by the TMA technique and by very careful

thermal analysis. The mischmetal itself contains at least four RE elements, Ce, La, Nd and Pr and the alloying behaviour of La with Co is quite different from that of the other RE elements with Co. The La-Co system does not contain stable phases like 1:3 and 2:17. Hence a systematic study was undertaken to establish the phase relationships in the technologically important composition region at 900°C using X-ray, metallography and TMA extensively and the EPMA to a limited extent.

An estimation of the primary magnetic properties, magneto-crystalline anisotropy field ( $H_A$ ), saturation magnetisation ( $4\pi$   $M_S$ ) and the Curie temperature ( $T_C$ ) is essential to decide the suitability of a RE-Co phase for permanent magnet potential. The MM-Co-Fe and MM-Co-Cu phases were studied for their  $4\pi$   $M_S$  and  $T_C$  using a vibrating sample magnetometer.

In the process of magnet production the parameters to be studied are: (i) composition of the alloy, (ii) size and size distribution of the powder (iii) compacting method (iv) sintering temperature and time and (v) post-sintering annealing temperature and time. These parameters affect both the B<sub>r</sub> and it and hence the (BH)<sub>max</sub>. The selection of right combination of the above parameters is essential to maximise the permanent magnet properties. Obviously, a systematic study varying all the above parameters will be a lengthy process and may not be necessary. Composition, powder characteristics, sintering temperature and sintering time are the parameters investigated

in the present study, only MM-Co alloys were tried in this connection. The powder production and the sintering are the most important steps in the magnet processing as the RE-Co alloy powders are highly susceptible to oxidation. The quality of the powder produced was checked by measuring their iHc rather than their average particle size since no single available technique has been found to give correct estimation of the particle size of the micron-size RE-Co magnetic powders.

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## 3. EXPERIMENTAL

The experimental methods followed in the present study are grouped into four sections, 1) alloy preparation, 2) phase analysis, 3) characterisation of magnetic properties and 4) magnet processing.

## 3.1 Alloy Preparation:

## 3.1.1 Raw Materials:

Indian mischmetal used in the study has the composition (analysed by x-ray fluorescence and complexometric methods<sup>1</sup>)

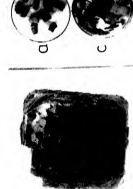
52 pct. Ce, 20.1 pct. La, 15.7 pct. Nd, 4.8 pct. Pr and 6 pct.

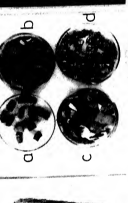
Fe (all in w/o) and the balance (~1.4 pct.) unidentified impurities. The Co, Fe and Cu used are of \$99.9 pct. purity.

The specifications of these materials are given in appendix I.

## 3.1.2 Melting:

The mischmetal ingot (Fig. 3.1) was cut into small pieces and ground to remove the outer oxidised surface. An effective gm. at. wt. of 140.06 gms. was computed for MM based on chemical analysis. The component elements MM, Co, Fe or Cu were weighed in the required proportions. The computation procedure of alloy composition is given in appendix II. The allowere prepared by arc melting in a water cooled Cu crucible in an ultra high purity argon atmosphere. Three or four meltings each lasting for 20 to 30 seconds were given and the alloy button weighing about 30 gms. was broken to small pieces and placed inverted after each melting to ensure uniformity in composition through out the ingot. The weight loss during









alloy button (4) Arc melted

## (a) Cut and polished MM (b) Co (c) Fe and (d) Cu Mischmetall Indian (2) Raw materials: Ingot

## 3 Arc melting unit

# ANALYSIS: EXPERIMENTAL PHASE







6 Annealing furnace set-up



(2) Homogenised alloy pieces

## PHASE ANALYSIS

- 2. X-ray diffraction 3. Thermomagnetic analysis (T.M.A. 1. Microstructure 4. Electron probe
  - micro-analysis (E.P.M.A.)

Materials, equipments, and methods employed in phase analysis melting due to evaporation of the RE elements was typically of 0.5 to 1 w/c.

## 3.1.3 Annealing

As the molten alloy is cooled very fast and most of the phases in RE-Co systems form through peritectic reactions segregation is inevitable. To homogenise the alloys small lumps of the as-cast alloy buttons were sealed in evaluated (10) quartz ampoules (Figure 3.1), annealed at 900°C for 4 to 10 days and quenched in water at the end of the annealing to retain the equilibrium phases corresponding to the annealing temperature. These homogenised alloy pieces were used for phase analysis.

## 3.1.4 Chemical Analysis

In arc melting of the alloys, the loss of RE due to evaporation is unavoidable and it varies depending on the arc pressure, melting time, amount of the alloy melted etc.

Hence it was necessary to check the composition of the alloys melted. The alloys were analysed for total RE, Co, Fe and Cu. As the Fe caused interference in the determination of RE and Co, it was precipitated quantitatively as Fe (111) hydroxide and analysed while the RE and Co were determined from the filtrate by visual complexometric method. The Cu in the alloys were determined either by complexometric method or by iodimetry. The detailed procedure of the chemical analysis is reported in Reference 1.

## 3.2 Phase Analysis:

Four techniques, metallography, x-ray diffraction, thermomagnetic analysis (TMA) and electron probe microanalysis (EPMA) were employed to study the phases.

## 3.2.1 Metallography:

A small piece of the homogenised alloy was taken and its surface for microstructural observation was prepared by standard procedures 2,3. The brittle alloys were hand ground carefully to avoid the introduction of cracks and then polished with fine alumina (0.l\u03c4) powder. The polished surfaces were then etched with 2 pct. nital solution for 2 to 3 seconds and the microstructure was observed with a Carl Zeiss NU-2 universal microscope in bright field.

## 3.2.2 X-ray Diffraction:

varied out with G.E. XRD-5 and 6 and a Philips PW 1730/10 x-ray diffractometers. For MM-Co-Fe alloys Cr, Co or Fe  $K_{\alpha}$  radiation was used but for high Cu containing alloys Cu  $K_{\alpha}$  radiation gave better results. Because of the close structural similarities 4-6 of the different RE-Co phases, their strongest reflections have close 'd' spacings and occur at close 20 values. In order to resolve them Cr  $K_{\alpha}$  radiation was used as the preferred one. powder samples were prepared by crushing the brittle alloys to -325 mesh size. The high Co and high Cu containing alloys were less brittle and tougher than the less Co or Cu bearing alloys. Hence Co or Cu rich alloy powders were annealed at  $600^{\circ}$ C for 10 mimutes to remove the strain prior to x-ray diffraction.

diffractograms were obtained by scanning at a rate of 2°/min.

In those cases where the lattice parameters of the phases were to be determined, the peaks were scanned at a slower speed (0.2°/min).

## 3.2.3 Thermomagnetic Analysis (TMA)

TMA was found particularly necessary to study the low cobalt phases with stoichiometry lower than 1:5 in the RE-Co systems (RE=Ce. La. Nd and Pr). The melting points of the peritectic 2:7 and 5:19 phases in these systems differ by less than 10°C and their standard x-ray diffraction data calculated based on their lattice parameters have the strong lines occurring at nearly the same position leaving only a few weak lines characteristic of these phases 7,8. Hence with conventional techniq such as thermal analysis and x-ray diffraction commonly used in determining phase diagrams, possibilities exist to misinterpret Also, when the phases 5:19, 1:5 and 2:17 coexist in a non-equilibrium condition in a case such as the partially decomposed 1:5 alloy, then the identification of 1:5 by x-ray diffis not possible as all the strong lines of 1:5 are common either to 5:19 or 2:17 phases 8-10. Since these phases are ferromagnetic and have sufficiently different Curie temperature (T<sub>o</sub>) 11, TMA was used to identify the phases and to obtain evidence for both the decomposition products of MMCo5. was used as a general tool like metallography or x-ray diffraction for phase identification in the present study. The experimental procedure is described under section 3.3.3.

of the area of the same of the

## 3.2.4 Electron Probe Microanalysis:

A model microsonde Electronic-MS-46 electron probe microanalyser was used to analyse the samples with an accelerating voltage of 20 KV and an x-ray emergence angle of 18°. The specimens were mounted in a perspex mold and their surfaces were prepared as for optical metallography. The polished surfaces were faintly etched with 1 pct. nital solution just to reveal the phase boundaries. This facilitated to select the right site for microanalysis of the elements. The phases were analysed for the five RE elements (Ce, La, Nd, Pr and Sm) and the three 3d transition elements (Fe, Co and Ni). Metals of the above elements with purity 99.9 pct. were mounted separately in different sample holders and used as standards to compare the x-ray intensities emitted from the samples. Both point and line scanes cutting across many grains were carried out.

3.3 Characterisation of Magnetic Properties:

## 3.3.1 Primary Magnetic Properties:

The three primary magnetic properties that determine the suitability of a RE-Co phase for permanent magnets are 1) the magnetocrystalline anisotropy field  $(H_A)$ , 2) saturation magnetisation  $(4\pi M_S)$  and Curie temperature  $(T_C)^{12,13}$ .

## 3.3.1.1 Magnetocrystalline anisotropy field, HA:

The  $H_A$  is experimentally determined by tracing the magnetisation curves along the magnetic easy and hard directions of a single crystal. The magnetic field corresponding to the point of intersection of the two curves gives the  $H_A^{12}$ . In the

case of polycrystalline material the same is obtained using a magnetically aligned powder specimen in which each powder particle is assumed to be a single grain. In practice the hard-axis magnetisation curve is plotted only upto \$\infty\$ 100 KOe owing to the limitation of the field and then extrapolated as the H<sub>A</sub> of the RECo<sub>5</sub> phases is much higher than 100 KOe. This gives in many cases unrealistic values. Hence the H<sub>A</sub> values estimated for oriented powder specimens in low magnetic field have to be treated with caution as they can give only an idea of the magnitude of the value. In the present study because of the non-availability of magnetic field greater than 11 KOe, the estimation of H<sub>A</sub> was not attempted for MM-Co intermetallic phases.

## 3.3.1.2 Saturation magnetisation, $4\pi M_s$ :

A PAR Model 155 vibrating sample magnetometer was used to estimate the specific magnetisation, σ (emu/gm) at ambient temperature. The magnetometer was calibrated with a standard PAR Ni sample which has a σ of 55 emu/gm at 298°K. All the magnetisation results reported in this work are based on this value. As the MM-Co phases are expected to have H<sub>A</sub> 10 KOe the alloy specimens were ground to powders of a few micron size under toluene. The powder was mixed into a paste with an epoxy resin and was filled in glass ampoules. Then they were aligned in a magnetic field (11 KOe) for few minutes and allowed to set in air in the absence of the field as they took 10 to 15 hrs for setting. The resin to powder weight ratio was typically of 0.5, the powder weight in the samples being in the range of 0.06 to 0.1 gm. Samples thus prepared have M<sub>r</sub>/M<sub>s</sub> ratio of

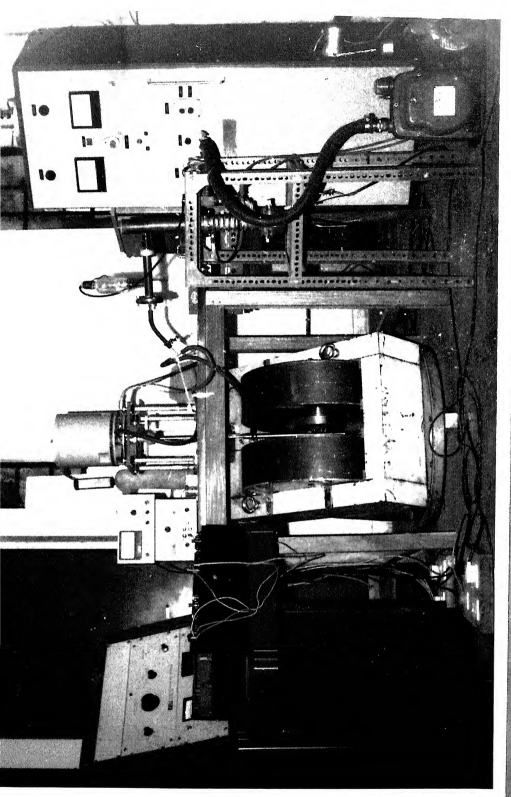
about 90 pct. and can be saturated with a 11 KOe field. When the magnetisation is reported for stoichiometric MM-Co phases, annealed samples were used in which the presence of the second phase as estimated by optical microscopy was \$\leq 5\$ pct.

3.3.1.3 Curie Temperature, Te:

ot studied.

As-cast as well as annealed alloys were used for thermomagnetic analysis (TMA). The samples studied were of about 0.2 gm. The TMA was carried out by measuring the magnetic moment (emu) of the sample as a function of temperature in a low magnetic field (40 0e). A PAR model 155 vibrating sample magnetometer coupled with model 151 high temperature oven assembly was used (Fig. 3.2). The alignment of the oven assembly was checked by measuring the T<sub>C</sub> of the standard PAR Ni sample which had a value of 356°C. The samples were taken in a boron nitride crucible and cept in a vacuum of 10<sup>-3</sup> Torr during the test to avoid oxidation. The output terminals of the Cr-Al thermocouple and the magnetometer were connected to the x and y axis, respectively, of an x-y recorder and the magnetic moment of the sample was continuously traced as a function of temperature. As the upper limiting temperature of the oven was 727°C, the phases having T<sub>C</sub> greater than 700°C were

Since the thermocouple of the PAR high-temperature oven ndicates only the oven temperature and the sample takes some ime to equilibriate with the changing oven temperature, a the hermal hysteresis was observed between the heating and cooling urves which decreased with decreased rate of heating or cooling.



8 pole gap, 2 air gap and 11 KOe peak field)(4) Vacuum systemhigh-temperature oven assembly (3) Polytronic electromagnet Fig. 3.2 Experimental set-up for thermomagnetic analysis (1) FAR model 155 vibrating sample magnetometer (2) Model 151

At a rate of 2°C/min a thermal lag of 10-15°C was persistent. the Hence both the heating and cooling curves were traced at the sar rate, especially near the magnetic transition regions and the mean of the transition temperatures read from these two curves was taken as the ferromagnetic Curie temperature of the phases studied.

## 3.3.2 Permanent magnet properties:

The permanent magnet properties of technical importance are Br, iHe, bHe and (BH) These parameters are obtained from the demagnetisation curve (II quadrant of the magnetic hysteresis loop) of the magnet samples. The experimental procedure to evaluate these properties is described under section 3.4.6.

## 3.4 Magnet Processing:

The magnet processing involves the following steps, alloy preparation, comminution, compacting, sintering, magnetisin and characterisation of the magnets for their physical and magnetic properties (Figure 3.3).

## 3.4.1 Alloy Preparation:

MM-Co alloys were prepared by arc melting method described in section 3.1. Each melt was about 30 to 40 gms. The as-cast arc melted buttons were used as such for powder production.











(C) Pot mill

(2) Crusher, sieve (-100 mesh) and brush

DArc melted alloy

button

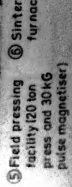
3 Stainless steel grinding jar and rods

# PROCESSING: EXPERIMENTAL MAGNET





(9) Magnetic characterisation: vibrating sample magnet meter and electromagnet



furnace set-up 6 Sintering

Materials, equipments, and methods employed processing in magnet

#### 3.4.2 Comminution:

The comminution of the as-cast alloys was carried out in two stages. First the alloy was hand crushed in a steel crusher to -100 mesh size in air and sieved into a small stainle jar (volume of 125 cc) containing toluene. Fine grinding was carried out by rod milling the -100 mesh powder in the stainless steel jar with stainless steel rods as the grinding media under sodium gettered toluene. The rods were of 2 mm dia and 9.5 cm long; the jar was of 4 cm dia and 10 cms in length. After milling, the wet powder was dried in a dessicator under vacuum (500µ). The optimum parameters to be selected in grinding are the feed size, feed volume and the milling time. In the present the feed size was fixed at -100 mesh as feeds finer than this may lead to severe oxidation of the powder. The effectiveness of the grinding and the optimum parameters of feed volume and milling time were determined by measuring the H of the powder rather than their average particle size. A feed volume of 5 gms was used. in the present study.

#### 3.4.3 Compaction:

The powders were field-pressed in a pulse magnetic fiel of 13 KOe applying a load of 14 Tons/cm<sup>2</sup>. A 20 Ton Bemco Hydraulic press and an RFL model 747-6 magnet charger fitted with a charging fixture of 3½ i.d. and 2 length was used (Fig. 3.3). The direction of the field and the pressing was parallel. Pellets of 0.96 cm dia and 0.3 to 0.6 cm thickness were formed wi a green density of 5.6 gm/cm<sup>3</sup> which is about 65 pct. of the

theoretical density for MM-Co phases. The powder compacts

were sintered immediately after pressing.

#### 3.4.4 Sintering:

The sintering of the powder compacts was carried out in a flowing argon (ultra high pure - 10LAR Grade) atmosphere at temperatures between 970°C and 1050°C in an alumina crucible. The sintering set up is shown in Fig. 3.3. One end closed alumi tube of 2.5 cm in i.d. and ~45 cms in length was fitted with a brass flange with an inlet and an outlet gas-leads of zirconia tubes of 3 mm bore. An alumina crucible containing the green pellets was placed inside the tube assembly and the end was closed vacuum tight. The chamber was evacuated to 20µ pressure and then flushed with argon. This was repeated 2 or 3 times. Then the alumina tube assembly was lowered into a furnace kept at any desired sintering temperature. It normally takes 10 to 15 minutes for the pellets to reach the sintering temperature. Sintering time was counted from the time the set sintering temperature was reached. At the end of the sintering period the alumina tube assembly was taken out and cooled in air. The cooling takes less than 3 min to reach 500°C and about 30 minutes to come to room temperature.

#### 3.4.5 Magnetising:

The sintered pellets were magnetised prior to magnetic measurements in an RFL model 747-6 magnet charger fitted with a charging fixture of l\* i.d. and 4" length. This can give a pulse field of 60 KOe with a pulse width of 100 to 200 milliseconds. The field was applied in a direction parallel to the pressing direction.

#### 3.4.6 Characterisation:

The permanent magnet properties  $B_r$ ,  $i^H c$ ,  $b^H c$  and  $(BH)_{max}$  of the sintered pellets were measured and compared with the density and phase and chemical composition of the pellets studied.

#### 3.4.6.1 Permanent magnet properties:

The B<sub>r</sub> and  $_{i}$ H<sub>c</sub> are obtained directly from the intrinsic demagnetisation curve (4 $\pi$ M vs. H) while the  $_{b}$ H<sub>c</sub> and (EH)<sub>max</sub> are obtained from the induction (B vs. H) demagnetisation curve. The latter curve is derived from the former using the relation B = H + 4 $\pi$ M where H is a negative quantity (demagnetising field).

A PAR model 155 vibrating sample magnetometer (VSM) coupled with a Polytronic type RTB-200 electromagnet which can give a peak field of 11 KOe, was used. The field was measured with a Hall probe - Gaussmeter. The output terminals of the Gaussmeter and the VSM were connected to an x-y recorder and the I and II quadrants of the hysteresis loop of the sintered and magnetised pellets were traced.

#### 3.4.6.2 Density:

The density of the sintered pellets was determined by geometrical as well as liquid (toluene) displacement methods.

#### 3.4.6.3 Phase composition:

To relate the magnetic properties observed to the structure of the sintered pellets, metallography and x-ray diffraction were performed. The techniques are the same as described under section 3.2. For x-ray diffraction the smooth sintered

pellet surface was used as such with no necessity of powder preparation.

### 3.4.6.4 Chemical composition:

Chemical analysis was carried out for a given alloy composition at three stages, 1) as-cast alloy, 2) powder compact and iii) sintered pellet. This was necessary to check any evaporation loss or oxidation of rare earth during the course of magnet processing. The procedure is the same as given under section 3.1.4.

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## 4. PHASE RELATIONS IN MM-Co-Fe SYSTEM IN CO-RICH REGION

#### 4.1 Introduction

Intermetallic phases of 1:5 and 2:17 stoichiometry in binary RE-Co (RE = Ce, La, Nd and Pr) systems have attractive magnetic properties: magnetocrystalline anisotropy field (H,), saturation magnetisation ( $4\pi$  M<sub>s</sub>) and Curie temperature (T<sub>s</sub>). These phases are used for the production of high energy density permanent magnets.  $^{6-12}$  The properties,  $^{H}_{i}$  and  $^{B}_{r}$  of these magnets are very sensitive to phase composition. 13-16 Hence a knowledge of phase relations near the 1:5 and 2:17 stoichiometry becomes important for selecting the right composition that gives a single phase alloy and for knowing how a slight change in composition alters the microstructure of these alloys. Although details of phase relations near 1:5 and 2:17 stoichiometries are available for pure binary RE-Co systems 17-20 phase relationships in a multicomponent MM-Co-Fe system are not investigated in detail. Mischmetal itself contains at least four RE elements (Ce. La. Nd and Pr). While the binary RE-Co phase diagram for RE = Ce, La and Nd are similar the La-Co system differs from the rest. It does not contain 1:3 and 2:17 phases. 21 A phase diagram for MM-Co system 22 and detailed phase relationships in the Co-rich regions of the MM-Co system 23 have been reported previously.

The Indian mischmetal also contains about 6 w/o Fe as impurity. The effects of Fe addition on the magnetic properties of MM-Co and RE-Co phases have been studied  $^{24-25}$ 

but no systematic study on the phase equilibria of RE-Co-Fe and MM-Co-Fe systems have been reported. Also the 1:5 and 2:17 phases in binary systems have been reported to have homogeneity region 17,19,20 and the former phase (1:5) undergoes an eutectoid reaction at low temperature (700°C). 29 Hence a systematic study was undertaken (i) to identify the different phases occurring in the MM-Co-Fe system (ii) to obtain their crystallographic details (iii) to establish the 900°C isothermal section of the MM-Co-Fe phase diagram (iv) to find the homogeneity region of the phases and (vi) to confirm the phase stability of the MMCo<sub>5</sub> phase.

For the purpose of drawing the phase diagram all the RE elements in mischmetal were assumed to constitute as a single component as far as their effect on phase-formation is concerned. However, we ther this assumption leads to any violation of the phase rule was considered carefully in this study.

#### 4.2 Experimental

The alloys were prepared using Indian mischmetal and 99.9% pure Co and Fe. The composition of mischmetal is the same as given under Section 3.1.1. The melting and annealing procedure of the alloys are described under Sections 3.1.2 and 3.1.3. The materials used and the methods followed in the phase analysis are illustrated in Figure 3.1 (Chapter 3). The alloys homogenised at 900°C

for 4 to 10 days were taken for phase analysis. The alloys studied for the stability of MMCo<sub>5</sub> were annealed at 700°C for 15 to 30 days. The phases were studied by metallography, X-ray diffraction, thermomagnetic analysis (TMA) and electron-probe micro-analysis. The experimental procedure is given under Section 3.2.

#### 4.3 Results and Discussion

The chemical and phase composition of the MM-Co-Fe alloys studied are given in Table 4.1 and the 900°C isothermal section of the MM-Co-Fe phase diagram is given in Figure 4.1. Excepting the D-group alloys, the alloy compositions plotted in the MM-Co-Fe phase diagram were analysed by TMA and by the conventional metallography and X-ray diffraction techniques. Six phases, 1:3, 2:7, 5:19, 1:5, 2:17 and B-Co were identified between 75 and 95 a/o Fe (Figure 4.1) close to the MM-Co binary line. In addition to them, a new phase designated as X, with a T<sub>C</sub> of about 340°C was present between 5:19 and 1:5 stoichiometries. The 2:7, 5:19, 1:5 and 2:17 phases were found to extend to considerable extent into the MM-Co-Fe ternary. At greater than 15 a/o Fe an S phase has been reported 30 to exist on the RE-rich side of 2:17 stoichiometry.

4.3.1 Identity and Crystallographic Details of Different Phases in the MM-Co-Fe System

# 4.3.1.1 2:7 and 5:19 Phases The alloy A-1 (Table 4.1) with composition on the

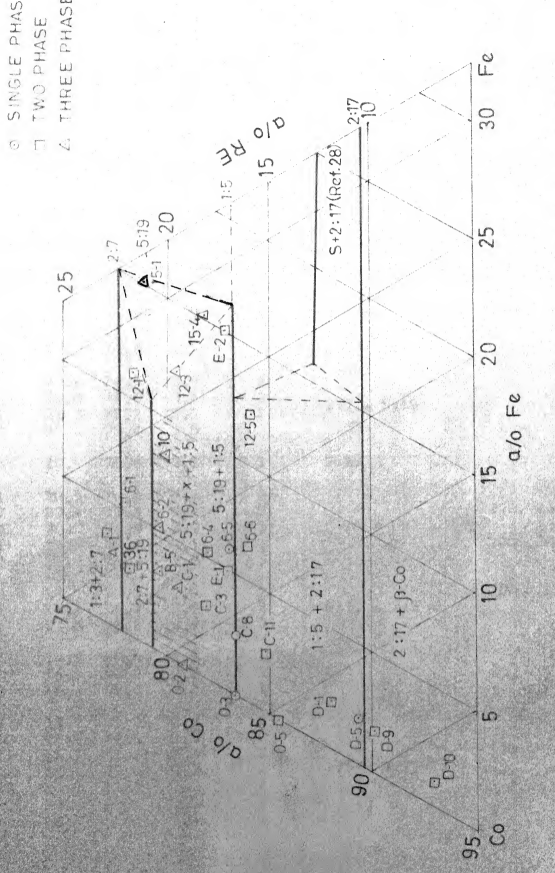


FIG. 4:1 900 °C ISOTHERMAL SECTION OF MM - CO - Fe PHASE DIAGRAM

TABLE 4.1 CHEMICAL AND PHASE COMPOSITION OF ALLOYS SELECTED FOR THE STUDY OF PHASE RELATIONS IN MM\_CO-Fe SYSTEM

Alloy		positi , anal		Pł	Phases (900°C, 4-10			
	MM <sup>a</sup>	Co	Fe	Micro- structure (No. of phases)	X-ray diffraction	TMAd (T <sub>C</sub> , °C)		
A-1 A-2	22.8 22.6	73.4 73.7	3.8 3.7	2, 2	1:3 + 2:7	<b>∠</b> 20, <u>6</u> 7		
A-3	22.3	74.0	3.7	1	2:7			
A-4 A-5 B-1 B-2	22.6 21.8 21.3 21.1	74.3 74.6 75.2 75.4	3.7 3.6 3.5 3.5	2 2 2 2	2:7 + 5:19			
B-3	20.9	75.6	3,5	1	5:19			
B-4 B-5 C-0 C-1 C-2	20.6 20.4 20.0 19.5 18.8	76.0 76.4 76.8 77.4 78.2	3.4 3.2 3.2 3.1 3.0	3 3 3 3 3	5:19 + (X) + 1:5	270 340 520 370 340 520 280 340 530		
C-3 C-4 C-5	18.1 17.4 17.5	79.0 79.7 79.6	2.9 2.9 2.9	2 2 2	5:19 + 1:5	280 480		
C-6 C-7 C-8	17.2 16.9 16.4	80.0 80.3 81.0	2.8 2.8 2.6	1 1 1	1:5			
C-9 C-10 C-11 C-12 C-13 C-14 C-15	16.2 15.9 15.1 14.6 14.1 13.6 13.2	81.2 81.6 82.5 83.1 83.6 84.1 84.6	2.6 2.5 2.4 2.3 2.3 2.3 2.2	2 2 2 2 2 2 2 2	1:5 + 2:17	520 <b>&gt;</b> 700		

Contd ...

Alloy	Composition (a/o, analysed)			P}	nases (900°C, 4-10	
	мм <sup>а</sup>	Co	Fe	Micro- structure (No. of phases)	X-ray diffraction	TMA <sup>d</sup> (T <sub>C</sub> , °C)
D-1 D-2 D-3 D-4	11.6 11.4	86.2 86.5 86.8 87.1	1.9 1.9 1.8 1.8	2	1:5 + 2:17	
D-5	11.0	87.2	1.8	1	2:17	
D=6 D=7 D=8 D=9 D=10	10.8 10.5	87.4 87.5 87.8 88.4 92.0		2	2:17 + β-Co	· ×
E-1 <sup>b</sup>	16.7	78.3	5,.0	2	5:19 + 1:5	280 550
E-2b	16.6	68.2	15.2	2	5:19 + 1:5	220 600
O-1C	19.4	80.6	0	3	5:19 + (x) + 1:5	
0-2°	19.2	80.8	0	3	5:19 + (X) + 1:5	<b>23</b> 0 <b>\$</b> 00 490
0-3 <sup>C</sup>	16.6	83.4	0	1	1:5	500
0-4°	15.8	84.2	0	2	1:5 + 2:17	0
0-5 <sup>C</sup>	14.6	85.4	0	2	1:5 + 2:17	
4-1	20.6	75.0	4.4	. 3	5:19 + (X) + 1:5	
4-2	20.2	75.6	4.2	3	5:19 + (X) + 1:5	
4-3	18.8	76.8	4.4	2 3 5 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	5:19 + (X) + 1:5	
4-4	17.6	77.9	4.5	3	5:19 + (x) + 1:5	
4-5	16.6	79.1	4.3	1	1:5	
4-6	15.9	79.9	4.2	2	1:5 + 2:17	Contd

Alloy	Composition (a/o, analysed)			Phases (900°C, 4-10 d)					
	MM <sup>a</sup>	Со	Fe	Micro- structure (No. of phases)	X-ray diffraction	TM.	a <sup>d</sup> c'°C	)	
6-1 6-2 6-3 6-4 6-5	22.3 20.5 19.5 18.0 16.6	75.3 76.7 77.0	5.1 5.2 5.3 6.4	2 3 3 2	2:7 + 5:19 2:7 + (X) + 1:5 2:7 + (X) + 1:5 5:19 + 1:5 1:5	130 270 260	320	2½0 540 525 530	
6-6 12-1 12-3 12-5 15-1 15-4	16.2 21.3 19.4 15.8 21.5 18.0	77.3 67.3 68.3 71.9 63.3 66.7	6.5 11.4 12.3 12.3 15.2 15.3	2 3 3 2 <b>3</b> 3 3	1:5 + 2:17 2:7 + 5:19 + 1:5 2:7 + 5:19 + 1:5 1:5 + 2:17 2:7 + 1:5! +5:19? 2:7 + 5:19 + 1:5	170 200	280 290 580 290	>600 700 >600 >600	

- a: MM represents only the total RE (Ce, La, Nd and Pr) content and does not include the Fe and the impurities present (1.4 w/o) in mischmetal. The Fe present as impurity in MM plus the Fe added intentionally is shown in a separate column. The analysed compositions (w/o) of the alloys for RE, Co and Fe were normalised to 100% and then converted to a/o. The average at. wt. of MM was taken to be 140.82 based on the X-ray fluorescence analysis.31
- b: E-1 and E-2 were prepared with synthetic MM of composition: 49.4% Ce, 22.8% La, 19.5% Nd, 4.9% Pr and 3.4% Sm (all in w/o).
- c: Alloys 0-1 to 0-5 were prepared with synthetic MM of composition: 48.4% Ce, 22.6% La and 29.0% Pr (all in w/o).
- d: The Curie temperatures shown in this table are indicative only for the purpose of identification of phases and the actual T<sub>C</sub> may differ by ±10°C.

rare-earth-rich side of the stoichiometry 2:7 shows two phases (Figure 4.2) and the alloy A-3 is a single phase. In the B-group, the alloy B-5 on the cobalt-rich side of the stoichiometry 5:19 has three phases and the alloy B-3 is a single phase. The X-ray diffraction data of the alloys A-3 and B-3 are given in Tables 4.2 and 4.3. Interestingly, both the alloys have all their strong lines occurring at nearly the same 20 position, leaving only a few weak lines which are characteristic of the phases. Thus it was ambiguous to conclude wether the phase in B-3 was different from that in A-3. To investigate this further, the alloys A-1 and B-5 were studied by thermomagnetic analysis and their thermomagnetic curves are shown in Figure 4.3.

The alloy A-1 shows one magnetic transition near 60°C whereas the alloy B-5 has a major transition near 270°C, two weak transitions near 340°C and 520°C and none near 60°C. The magnetic transition observed at 60°C for the alloy A-1 is attributed to the MM<sub>2</sub>Co<sub>7</sub> phase (Figure 4.3). The magnetic transition of the second phase present in A-1 (MMCo<sub>3</sub>) is not observed as it should have a ferromagnetic transition temperature,  $T_{\rm C}$  much less than room temperature (20°C). Similarly the magnetic transition seen at 270°C for the alloy B-5 is attributed to MM<sub>5</sub>Co<sub>19</sub> phase and the other transitions are attributed to minor phases. These results confirm that the matrix phases in the alloys A-1 and B-5 are MM<sub>2</sub>Co<sub>7</sub> and MM<sub>5</sub>Co<sub>19</sub> respectively.

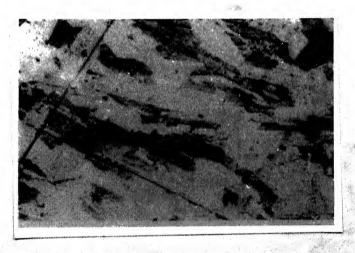


Figure 4.2 The microstructure of the alloy A-1. The areas are MMCo3 and the matrix is MM2Co7.

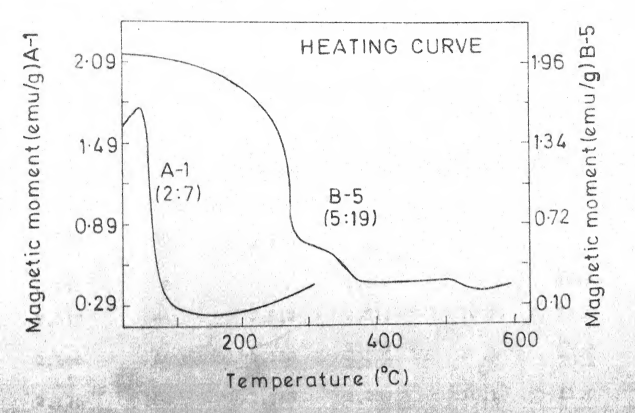


FIG. 43 THE LOW FIELD MAGNETIC MOMENT AS A FUNCTION OF TEMPERATURE FOR THE ALLOYS A-1 AND B-5 AFTER ANNEALING AT 900 °C FOR 10 DAYS

TABLE 4.2 X-RAY DIFFRACTION DATA FOR MM2Co7 PHASE

Hexagonal  $\underline{a}: 4.99_{0}$  Rhombohedral  $\underline{a}: 5.02_{4}$ 

<u>c</u>: 24.69<sub>9</sub>

c: 36.71<sub>3</sub>

d <sub>obs</sub> (A)	% <sup>I</sup> obs.	d cal.	( Å)	(hk	(1)
ous.	DDS.	hex.	rhomb.	hex.	rhomb.
2.804	11	es.	2.806	Ų a	10.1
2.745 2.724	18 20	2.733	ea	10.7	
2.645	8	==	2.648	==	10.11
2.513	61	2.512	2.512	10.80	11.00
2.166	46	2.153	2.172 2.160	20.1	20.1
2.136	100	2.144	2.139	10.1	11.9
2.034	23	grale	2.039	439	00.18
1.938	6	1.941	1.934	11.80	10.17
1.745	7	1.739	1.748	10.13	00.21 11.15
1.581	4	603	1.583	<b>4</b>	11.8
1.451	7	1.454	1.475	10.16	21.10
1.433	7	1.427	1.445	20.13	20.19 30.6
*1.367	35	1.367	1.367	20.14	30.9
*1.256	31	1.256	1.256	20.16	22.00

<sup>\*</sup> Lines taken for the calculation of lattice parameters.

<sup>-</sup> Denotes not present.

TABLE 4.3 X-RAY DIFFRACTION DATA FOR MM  $_5$ Co  $_{19}$  PHASES Rhombohedral - a: 5.01 $_8$ ;  $\underline{c}$ : 48.65 $_7$ 

	1		
d <sub>obs</sub> (Å)	% <sup>I</sup> obs.	d <sub>cal.</sub> (Å)	(hk.1)
2.831 2.798	9 16	2.836	10.13
2.714	13	2.714	10.14
2.509*	43	2.509 2.492	11.00 10.16
2.166	39	2.171 2.164	20.1% 20.20
2.134	100	2.134	11.2
2.027*	23	2.027	00.24
1.578	4	1.577	11.24
1.449	8	1.448	30.00
1.364	26	1.364	30.12
1.254	23	1.254	22.1

<sup>\*</sup> Lines taken for the calculation of lattice parameters.

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The X-ray diffraction lines of the alloy A-3 were indexed based on the hexagonal Th2Co7 type structure. 32 However, some of the lines could not be accounted for based on this structure. Khan $^{33}$  has reported that for  $\text{RE}_2\text{Co}_7$ (RE = Ce, Pr, Nd and CeMM) the rhombohedral modification (Gd2Co7 type) is the stable low temperature form whereas Buschow 32 has reported that both the modifications could coexist. In the present study the observed diffraction lines for the alloy A-3 could be accounted for only if both the modifications were assumed to be present (Table 4.2). The lattice parameters calculated for both the modifications are shown in Table 4.4. All the X-ray diffraction lines of the alloy B-3 could be indexed on the basis of the rhombohedral Ce<sub>5</sub>Co<sub>19</sub>-type structure 34 and the lattice parameters calculated for the MM<sub>5</sub>Co<sub>19</sub> phase in the alloy B-3 are in close agreement with the values reported for  $CeMM_5Co_{19}$  by Khan  $^{33}$ (Table 4.4).

Alloys 6-1, 12-1 and 15-1 (Figure 4.1) lie between the 2:7 and 5:19 stoichiometries and the Fe content increases from 5.4 a/o in alloy 6-1 to 15.2 a/o in alloy 15-1 (Table 4.1). Their microstructures show two phases. These alloys were also analysed by TMA and the Curie temperature of the phases (Table 4.1) show a gradual increase from 67°C in alloy A-1 to 200°C in alloy 15-1 for the 2:77 phase. Similarly for the 5:19 phase the T<sub>C</sub> was found to increase from 220°C in alloy 36<sup>35</sup> to 280°C in alloy 12-1. This increase

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TABLE 4.4 COMPARISON OF LATTICE PARAMETERS OF STOICHIOMETRIC PHASES OF MM\_Co SYSTEM WITH THOSE OF Ce\_Co AND CeMM-Co SYSTEMS

Phases	Structure	Crystal	Lattice parameters (Å)					
	type	symmetry	MM-Co		Ce_Co <sup>33,34</sup>		CeMM-Co <sup>33</sup>	
		f l l	a	C	a	C	<u>a</u>	c
2:7	Gd <sub>2</sub> Co <sub>7</sub>	rhomb.	5.024	36.713	4.956	36.5 <sub>25</sub>	5.023	36.747
	Ce <sub>2</sub> Ni <sub>7</sub>	hex.	4.99 <sub>0</sub>	24.699	4.944	24.470	ei3	-
5:19	Pr <sub>5</sub> Co <sub>19</sub>	rhomb.	5.018	<sup>48</sup> .6 <sub>57</sub>	4.947	48.743	5.012	48.716
1:5	CaCu <sub>5</sub>	hex.	4.98 <sub>O</sub>	4.008	4.922	4.030	4.991	4.004
2:17	Th <sub>2</sub> Zn <sub>17</sub>	rhomb.	8.472	12.145	8.378	12.206	c=	er.as

is attributed to the presence of Fe in these phases. The 2:7 phase does not exist in the RE-Fe binary system. The present study indicates the 2:7 phase extends into the termary as a stable phase at least upto 15 a/o Fe. The 5:19 phase was not observed in alloy 15-1 and the alloy 12-3 contains three phases. The 5:19 phase does not seem to extend to greater than 10 a/o Fe into the MM-Co-Fe termary.

#### 4.3.1.2 1:5 Phase

The alloy c-6 (Table 4.1) was found to be a single phase and its X-ray diffraction pattern could be indexed on the basis of the  $CeCo_5$ -type structure. The X-ray diffraction data of the MMCo<sub>5</sub> phase is given in Table 4.5. This

TABLE 4.5 X-RAY DIFFRACTION DATA FOR MMCo<sub>5</sub> PHASE

Hexagonal

<u>a</u>: 4.98<sub>0</sub> Å <u>c</u>: 4.00<sub>8</sub> Å

d <sub>obs.</sub> (Å)	% I <sub>obs.</sub>	d <sub>cal</sub> .(Å)	(h k•1)
4.007	3	4.008	0 0 .1
2.930	38	2.936	1 0.1
2.483	29	2.489	1 1.0
2.154	41	2.156	2 0.0
2.113	100	2.115	1 1.1
2.003	19	2.004	0 0.2
1.898	17	1.899	2 0.1
1.562	16	1.561	1 1.2
1.509	14	1.510	2 1.1
1.468*	21	1.468	2 0.2
1.437	5	1.438	3 0.0
1.352	24	1.353	3 0.1
1.245*	31	1.245	2 2.0
		A second second	

Lines taken for the calculation of lattice parameters.

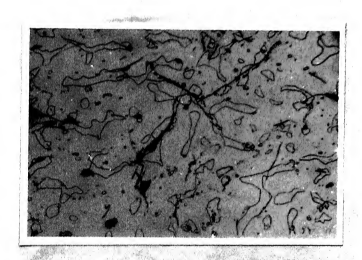


Figure 4.4 The microstructure of the alloy C-4. The islands are MM<sub>5</sub>Co<sub>19</sub> and the matrix is MMCo<sub>5</sub>.



Figure 4.5 Microstructure of alloy 6-6. The matrix is MMCo<sub>5</sub> and the minor phase is MM<sub>2</sub>Co<sub>17</sub>.



Figure 4.6 Microstructure of alloy 12-5. The matrix is MMCo<sub>5</sub> and the minor phase is MM<sub>2</sub>Co<sub>17</sub>.

phase appeared initially in the alloy B-4 and then increased in quantity with increasing cobalt concentration. The microstructure of the alloy c-4, which contains  $\text{MM}_5\text{Co}_{19}$ , and  $\text{MMCo}_5$  (matrix) is shown in Figure 4.4. The alloys c-6, c-7 and c-8 showed a single-phase (MMCo $_5$ ) structure (Table 4.1), indicating the presence of a homogeneity region for this phase. This is discussed further in the next section.

Alloy 0-3 was prepared with synthetic mischmetal and contains no Fe. This alloy shows single phase, 1:5 both in the microstructure and/the TMA. A number of alloys, c-8, 4-5, E-1, 6-5, 12-5 and E-2 (Figure 4.1 and Table 4.1) containing different amount of Fe along the 1:5 stoichiometric line were prepared. Alloys c-8, 4-5 and 6-5 show microstructurally single phase with a trace of second phase while the rest contain 1:5 as the predominant phase with small amount of the second phase. The second phase present in the alloy E-2 was the 5:19 phase as evidenced by its TMA. 38 The TMA pattern of the alloy 12-5 (Figure 5.6, Chapter 5) does not show any magnetic transition corresponding to the 5:19 phase. Hence the second phase seen in the microstructure of alloy 12-5 is attributed to 2:17 phase (Figure 4.5). Similarly the second phase seen in the alloy 6-6 (Figure 4.6) which gets etched brighter than the matrix 1:5 phase is attributed to the 2:17 phase. In earlier study the composition of 1:5 phase was reported to deviate towards higher Co content as the Fe content in the 1:5 increased. But the present study shows

no such deviation from the ideal stoichiometric line and the 1:5 phase exists as a stable phase at 900°C at least upto 17 a/o Fe in the MM-Co-Fe termary system.

#### 4.3.1.3 2:17 Phase

A phase having a crystal structure of the  ${\rm Th}_2{\rm Zn}_{17}$ type first appeared in the alloy C=9 and increased in quantity with increasing Co concentration (Table 4.1). A single phase of stoichiometry 2:17 was obtained for the alloy D-5, the X-ray diffraction data of which is given in Table 4.6. In between C-9 and D-5,  $\text{MMCo}_5$  and  $\text{MM}_2\text{Co}_{17}$  are present. The microstructure of the alloy C-11 in which the bright MM2Co17 is precipitated along the grain boundaries of  $\ensuremath{\mathsf{MMCo}}_5$  , is shown in Figure 4.7. The binary La-Co system does not have the La<sub>2</sub>Co<sub>17</sub> phase. However, in the present study it was found that natural mischmetal, which contains about 20 w/o La, forms a stable MM2Co17 phase and the observed X-ray diffraction lines for this phase could be indexed on the basis of the Th<sub>2</sub>Zn<sub>17</sub> rhombohedral structure. 39 This phase extends far into the ternary system and possibly upto the binary MM\_Fe.30

#### 4.3.1.4 β-Co Phase

In addition to the stoichiometric phases, a Co-Fe phase was found to coexist with MM2Co17 on the Co-rich side of 2:17 stoichiometry. Unlike the other alloys, the alloy D-10 (Figure 4.8), which contained about 40% of this

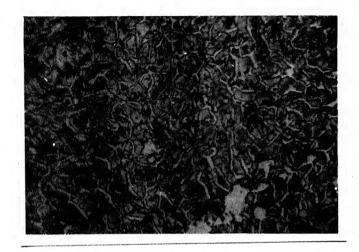


Figure 4.7 The microstructure of the alloy C-11. The matrix is  ${\rm MMCo}_5$  with  ${\rm MM}_2{\rm Co}_{17}$  at the grain boundaries.

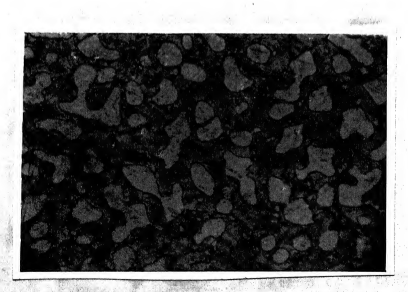


Figure 4.8 The microstructure of the alloy D-10. The matrix is  $MM_2^{Co}_{17}$  with islands of  $\beta$ -Co.

TABLE 4.6 X-RAY DIFFRACTION DATA FOR MM2Co17 PHASE

Rhombohedral  $\underline{a}$ : 8.47<sub>2</sub>  $\overset{\circ}{A}$ 

<u>c</u>: 12.14<sub>5</sub> Å

d <sub>obs</sub> .	% I obs.	$\mathtt{d}_{\mathtt{cal.}}^{}(\hat{\mathbf{\hat{\hat{\mathbf{A}}}}})}$	(hk.1)
3.138	4	3.139	20.20
2.929	41	2.926	11.30
2.837	8	2.806	10.40
2.689	8	2.703	21.10
2.513	9	2.523	21.20
2.431	36	2.446	30.00
2.347	34	2.339	20.40
2.102	60	2.118	22.00
2.085	100	2.093	31.30
2.042	41	2.047	21.40
2.031	44	2.024	00.60
1.870	31	1.877	22.30
1.833	8	1.826	11.60
1.804	5	1.814	40.10
1.745	5	1.737	22.40
1.611	6	1.606	11.70
1.562 1.480 1.476 1.464*	5 15 15 25	1.559 1.489 1.471 1.464	30.60 41.30 21.70 40.50 32.40
1.403 1.326 1.315 1.256	5 15 5	1.412 1.333 1.321 1.256	33.00 33.3 50.4 41.60
1.214*	31	1.214	00.10
1.209	8	1.222	60.0
1.203	6	1.217	60.1
1.200	8	1.200	10.10

<sup>\*</sup> Lines taken for the calculation of lattice parameters.

phase, was not brittle. The X-ray diffraction results confirmed that it existed as f.c.c.,  $\beta$ -Co. The  $\beta$ -Co phase was analysed using electron probe micro-analyser (EPMA) for the presence of any rare earth elements. Figure 4.9 shows an EPMA trace of  $\beta$ -Co and the MM<sub>2</sub>Co<sub>17</sub> phase of the alloy D-10; it was analysed for the three 3d elements, cobalt, iron and nickel, and for the five rare earth elements cerium, lanthanum, neodymium, praseodymium and samarium. For both the phases, except for Co, the elements were probed at the same X-ray intensity scale (300 counts per second). It can be seen that the  $\beta$ -Co phase contains predominantly Co with small amounts of Fe and Ni and contains no detectable amount of RE elements. The binary phase diagrams of the RE-Co systems 18 also show the absence of terminal solid solubility of rare earths in cobalt.

#### 4.3.1.5 X-phase

A phase having a T<sub>C</sub> about 340°C was observed for a number of alloys adjacent to 5:19 stoichiometric line on the Co rich side and the region where it was detected is shown in hatched form in the MM-Co-Fe phase diagram (Figure 4.1). This phase was initially suspected for the 1:4 type phase based on similar such evidence reported by Strnat for Pr-Co system. In order to confirm this MMCo<sub>4.0</sub> alloy (C-O) was prepared and its TMA curve was obtained which is shown in Figure 4.10. It is seen from this figure, that the

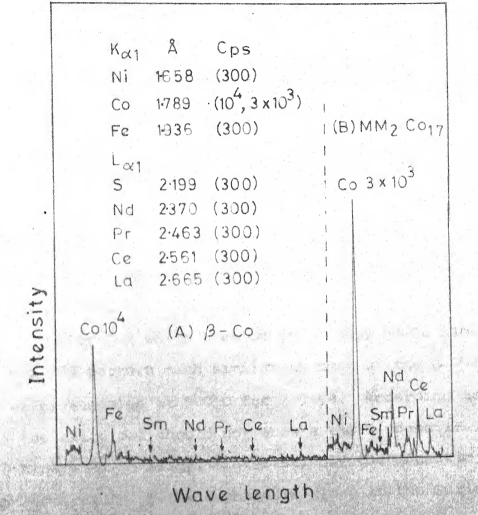
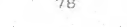


FIG. 4.9 AN EPMA TRACE OF THE ALLOY D-10
ANALYSED FOR 3d AND RARE EARTH
ELEMENTS IN 3-Co AND MM2 Co17

340°C transition still remains a minor one. It occurs in the 900°C annealed alloy and is absent in the as-cast and 1000°C annealed alloys (Figure 4.11). MMCo<sub>4.0</sub> alloy annealed at 700°C had similar TMA pattern as that of the 900°C annealed alloy. Also this phase was not present in alloy C-3, 6-4 and R-1 which are close to the 1:5 stoichiometric line (Figures 4.1 and 4.10) suggesting that this could not be due to 1:4 type of phases.

Alloy 0-2 which lies on the binary MM-Co line also shows a TMA pattern much similar to that of the C-O alloy for its allow annealed at 900°C for 7 days. According to phase rule the alloy 0-2 should not show a third phase if it were a true binary MM-Co alloy. Interestingly the magnetic transition temperature of the X-phase is close to the Curie temperature of the  ${\rm La_5Co}_{19}$  phase  $^{4\rm O}$  (343°C). A comparison of the peritectic temperatures of the different intermetallic phases in the RE-Co system for RE = Ce, La, Nd and Pr (Section 1.3.1, Chapter 1) indicates that for La5Co19 and La2Co7 the peritectic temperatures are lower by about 300°C from the other RE5Co19 and RE2Co7 phases. Wwether this will have any effect on phase separation of La5Co19 from the MM5Co19 is not known for certain. The X-ray diffraction results of MMCo4.0 did not reveal this phase possibly because the La5Co19 and MM<sub>5</sub>Co<sub>19</sub> have very close lattice parameters. However, in the microstructure an evidence could be obtained for the presence of small amount of a third phase. But wether it



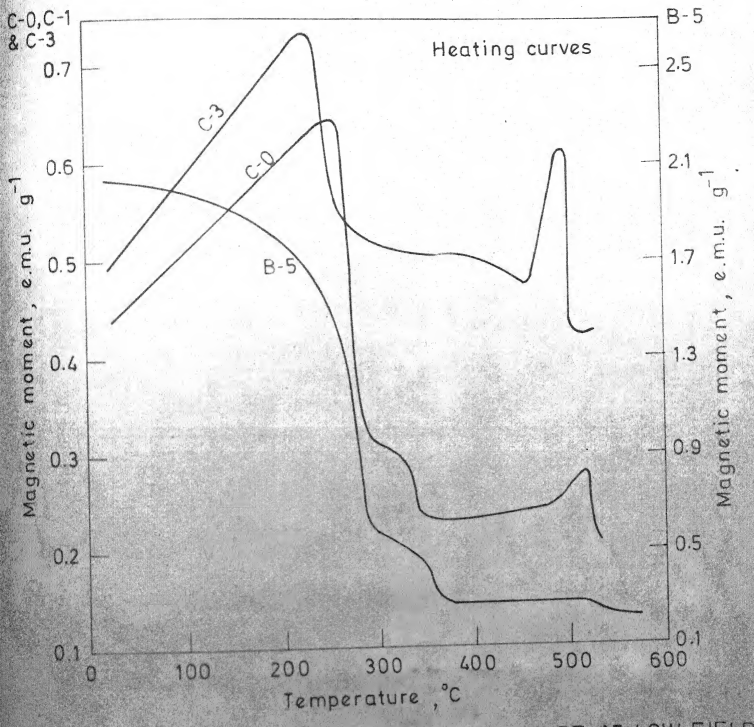


FIG 410 MAGNETISATION VS TEMPERATURE AT LOW FIELD FOR ALLOYS B-5, C-0 AND C-3, 900 C ANNEALED

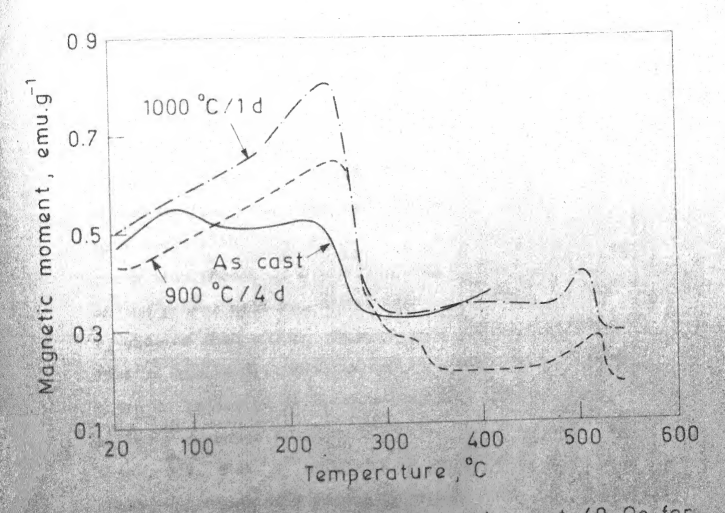
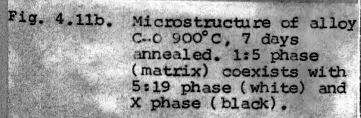


Fig.411 Magnetisation versus temperature at 40 Oe for MM Coza alloys (C-0)





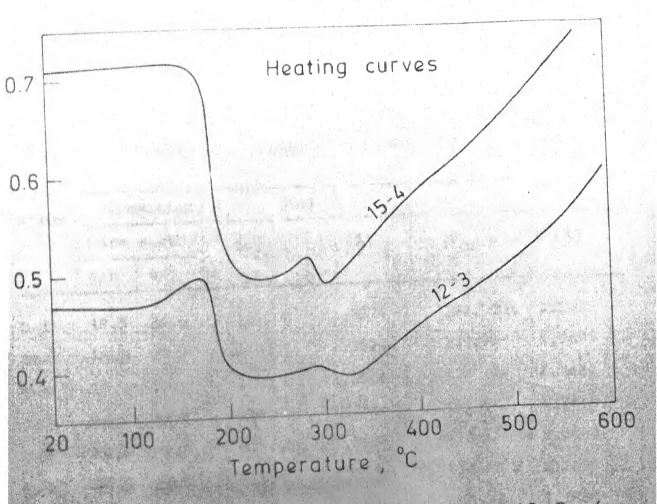
is really due to La<sub>5</sub>Co<sub>19</sub> is yet to be proved by using other techniques like EPMA.

# 4.3.1.6 High Fe Containing MM\_Co-Fe Alloys Between 2:7 and 1:5 Stoichiometries

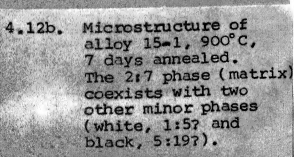
The phases of alloys present in this region all have closely related crystal structures and their X-ray diffraction results could not give unambiguous evidence for the identity of the phases. Hence TMA was used as a reliable technique to establish the identity of the phases. While the low Fe containing phases in this region had T less than 500°C, some of the phases with Fe content greater than about 10 a/o have To greater than  $600^{\circ}$  C. This is shown for alloys 12-3 and 15-4 in Figure 4.12 where a definite evidence for a third phase is indicated in their TMA patterns. However since 600°C is the safe upper limit of the high temperature oven used in this study, their T could not be determined. Alloy 15-1 shows phases only in the microstructure. Hence based on the microstructural results and the incomplete TMA results a tentative three-phase region is indicated in the MM-Co-Fe phase diagram.

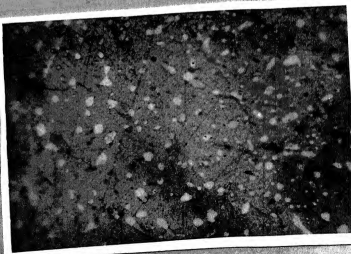
#### 4.3.2 Homogeneity Region of 1:5 Phase

In the study of the X-ray diffraction patterns of the alloys of near 1:5 stoichiometry, a systematic shift in the 20 position of the X-ray diffraction lines of the MMCo<sub>5</sub> phase was observed. The shift was distinctly visible on



412 Magnetisation versus temperature at 40 De for alloys 12 - 3 and 15 - 4; 900 °C, 7 days annealed





the high angle (20 > 100°) side and is given in Table 4.7 for the planes (30.1) and (22.0) of the MMCo<sub>5</sub> phase. The X-ray diffraction peaks of these planes were scanned at a slow rate (0.2°/min) in order to read the 20 value accurately ( $\pm$  0.02°).

TABLE 4.7 VARIATION IN THE POSITION (20) OF X-RAY DIFFRACTION LINES OF 1:5 PHASE WITH COMPOSITION

Alloy	lloy Composition  Rare earth		30.	1	22.0		
			<sup>20</sup> Cr	d (Å)	<sup>20</sup> Cr	d (Å)	
	a/o	w/o	Cr		CI		
C-1	19.5	36.5	115.00	1.3574	132.92	1.2487	
C-2	18.8	35.5	115.05	1.3570	132.95	1.2485	
C-4	17.4	33.6	115.10	1.3566	133.03	1.2482	
C-7	16.9	32.7	115.33	1.3549	133.37	1.2466	
C-8	16.4	31.8	115,45	1.3538	133.62	1.2454	
C-10	15.9	31.0	115.48	1.3537	133.66	1.2452	
C-11	15.1	29.8	115.52	1.3535	133.72	1.2450	
C-12	14.6	28.9	115.59	1.3530	133.78	1.2447	

The X-ray diffraction and microstructure results given in Table 4.1 also shows that the alloys C-6, C-7 and C-8 are single phase MMCo<sub>5</sub>. Using the reflections (30.1) and (22.0) (Table 4.7) the lattice parameters a and c of the MMCo<sub>5</sub> phase were calculated. The variation of the

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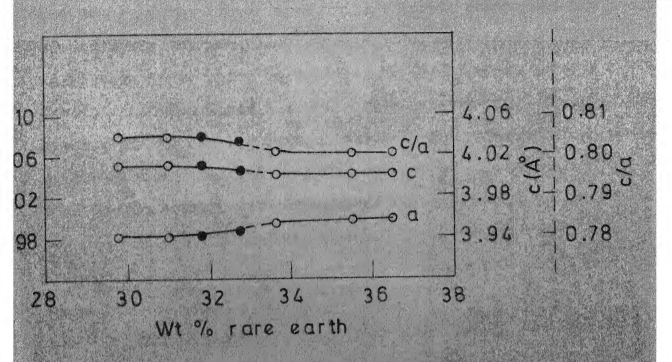
parameters <u>a</u> and <u>c</u> with composition is shown in Figure 4.13. In the single phase region, between alloys C-4 and C-10 (Table 4.7) <u>a</u> decreases and <u>c</u> increases (with a consequent increase of c/a) with increasing Co content. This is in agreement with the variation of lattice parameters reported by Buschow et.al. <sup>17</sup> and by Khan <sup>19</sup> in the homogeneity region of RECo<sub>5</sub> phase (RE = Sm, Ce, Pr and Nd). Khan <sup>19</sup> has reported the existence of a homogeneity region for  $CeCo_5$ ,  $PrCo_5$  and  $NdCo_5$  near  $1100^{\circ}C$ . The present study shows that mischmetal which is a mixture of Ce, La, Nd and Pr also has a homogeneity region for the MMCo<sub>5</sub> phase extending over 2 w/o (1.5 a/o) rare-earth at  $900^{\circ}C$  and its width is comparable with that reported by Khan <sup>19</sup> for the  $CeCo_5$  phase.

In the present study the homogeneity region was studied only for 1:5 phase alloys containing about 3 a/o Fe.

It was not studied closely for low as well as high Fe containing 1:5 alloys.

# 4.3.3 Phase Stability of MMCo5

The alloy C-6, which is single-phase MMCo<sub>5</sub> at 900°C, gave an interesting microstructure when annealed at 700°C for 30 days (Figure 4.14). A bright phase MM<sub>2</sub>Co<sub>17</sub> appeared along the grain boundaries of the MMCo<sub>5</sub> phase. Buschow  $^{27}$  has obtained a similar microstructure for the NdCo<sub>5</sub> phase annealed at 620°C for 8 weeks. He has attributed this to a eutectoid reaction in which the NdCo<sub>5</sub> phase decomposes into Nd<sub>5</sub>Co<sub>19</sub>



4-13 VARIATION IN THE LATTICE PARAMETERS WITH COMPOSITION OF THE MMCo<sub>5</sub> PHASE IN MMCo<sub>5</sub> ± x ALLOYS. •, Homogeneous alloys;
•, Heterogeneous alloys

and Nd<sub>2</sub>Co<sub>17</sub>, the latter appearing as bright grain boundary precipitates and the former appearing as dark parallel lines inside the NdCo<sub>5</sub> phase with different orientations in each grain. X-ray evidence has been given only for the Nd<sub>2</sub>Co<sub>7</sub> product phase. In the present study it was observed that the extent of decomposition of MMCo<sub>5</sub> even after annealing for 30 days was so small (<5%) that clear X-ray evidence could not be obtained for the product phases. Since the addition of iron to the phase of stoichiometry 1:5 has been reported to raise the eutectoid reaction temperature, <sup>41</sup> two alloys E-1 and E-2 (Table 4.1) containing 5 and 15.2 a/o Fe respectively, were prepared using synthetic mischmetal and high purity Fe and Co (>99.9%). The synthetic mischmetal contained high purity (>99.9%) rare earth elements in approximately the same ratio as that in natural mischmetal.

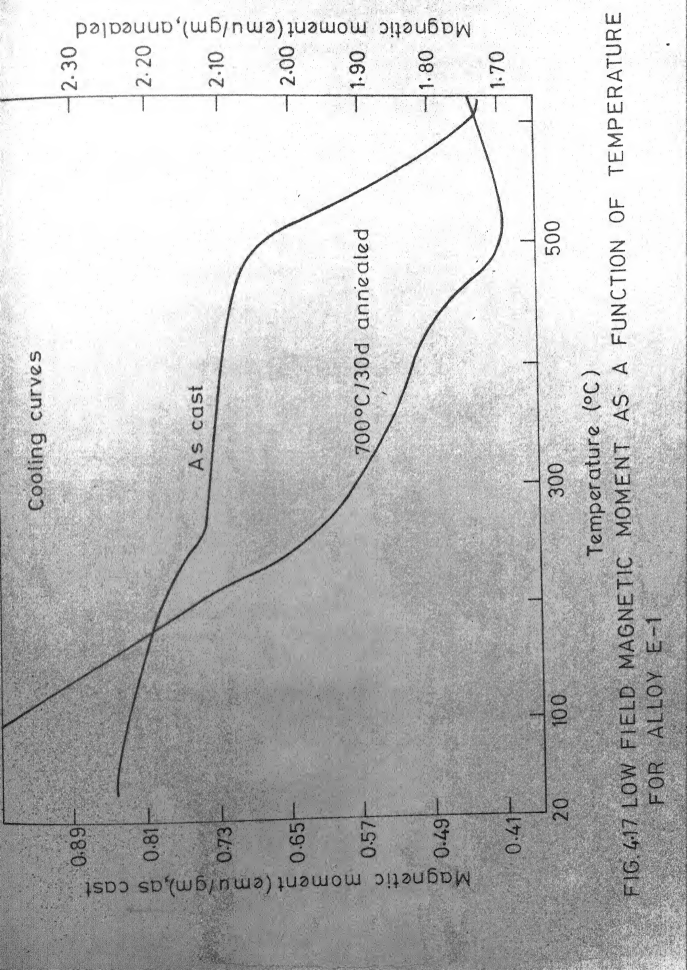
In Table 4.8 the X-ray diffraction results of as-cast and annealed alloys of E-1 and E-2 are shown. In the ascast condition both E-1 and E-2 contain predominantly  $^{\rm MMCo}_5$  with a small amount of  $^{\rm MM}_5{\rm Co}_{19}$ . The same alloys on annealing at 700°C for 30 days gave new reflections belonging to the  $^{\rm MM}_2{\rm Co}_{17}$  phase.

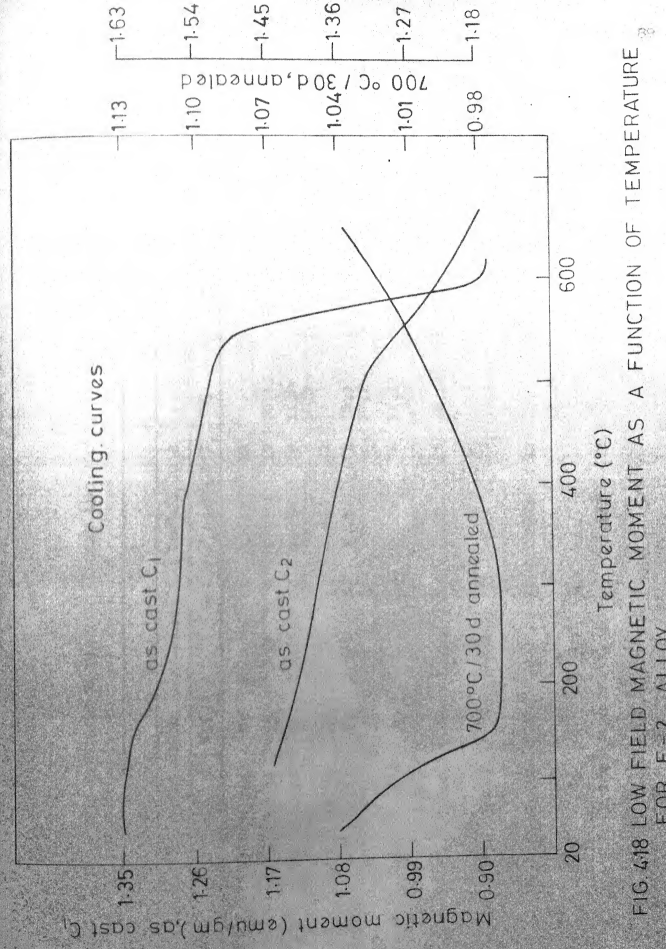
The microstructures of the alloys E-1 and E-2 annealed at 700°C for 30 days are shown in Figures 4.15 and 4.16. For the alloy E-1 the bright MM<sub>2</sub>Co<sub>17</sub> phase is precipitated along the grain boundaries and inside the grain (Figure 4.15) whereas in the alloy E-2 they appear as

separate grains (Figure 4.16). However, it was not possible to prove the absence of the parent  ${\rm MMCo}_5$  phase in these alloys because all the strong X-ray diffraction lines of  ${\rm MMCo}_5$  overlap with those of either the  ${\rm MM}_5{\rm Co}_{19}$  or the  ${\rm MM}_2{\rm Co}_{17}$  phase (Table 4.8). In the microstructure also no clear evidence could be obtained for this. Hence these alloys were further studied by thermomagnetic analysis.

In Figure 4.17 the TMA curves for the as-cast alloy of E-1 and for the alloy of E-1 annealed at 700°C for 30 days are shown. For the as-cast alloy a strong magnetic transition near 541°C and a weak magnetic transition at 240°C can be seen, belonging to MMCo<sub>5</sub> and MM<sub>5</sub>Co<sub>19</sub> respectively. On annealing, the MMCo<sub>5</sub> magnetic transition is greatly reduced and that of MM<sub>5</sub>Co<sub>19</sub> becomes prominent. The magnetic transition corresponding to the MM<sub>2</sub>Co<sub>17</sub> phase could not be seen in this curve because it has a higher ferromagnetic transition temperature, T<sub>C</sub> than the upper limit of the high temperature oven used. These results prove that the E-1 alloy annealed at 700°C contains mainly the MM<sub>5</sub>Co<sub>19</sub> and MM<sub>2</sub>Co<sub>17</sub> phase with a small amount of the parent MMCo<sub>5</sub> phase.

The TMA curves of the alloy E-2, which contains a greater amount of iron (15.2 a/o) than the alloy E-1 (5 a/o) show a different result (Figure 4.18). The MMCo<sub>5</sub>-magnetic transition observed for the as-cast alloy was totally absent for the alloy annealed at 700°C for 30 days, proving the absence of the MMCo<sub>5</sub> phase. The magnetic transition





ALLOY FOR E-2

TABLE 4.8 X-RAY DIFFRACTION DATA OF AS-CAST AND 700°C, 30 days ANNEALED ALLOYS OF E-1 AND E-2

	a.		lave .		Θ	E-2				hk.1 (% I <sub>C</sub> )	( I C)		
As-cast	ast	700°C, 30 d	30 d	As-cast	Ĭ.	700°C,	30 d				L		
d Oi	O T %	0 0 0	° 1 %	g	% I	່ວ	% I	ST:C	<u> </u>	T:2	۵	7:17	
2,942	4	2,942	33	2.942	38	2.942	28			10.1	(38)	11.3	(41)
		2,819	17	2,819	on	2.819	17	10,13	(16)			10.4	(8)
2,485	28	2,496	31	2,485	28	2.496		11.0	(43)	11,0	(29)		
		2,441	22	in the second		2,441	21	) •			,	30.0	(38)
2.149	<b>4</b> 6	2,157	41	2.141	47	2.137	72	20.1 20.2	(38)	20.0	(41)		
2.116	100	2,116	94	2,116	100	2.116	75	11.12	(100)	11.1	(100)	22.0	(09)
		2,093	100			2,101	100					31,3	(100)
		2.040	52	2,048	19	2,055	47	00.24	(23)			00.6	(44)
2,012	24	2,022	50	2.022	29					00.2	(13)		
1,893	14			1.893	13					20.1	(11)		
		1,869	17			1.875	19	,				22.3	(31)
1.559	12	1,559	6	1,583	0			11,24	(4)	11.2	(16)	30.6	(2)
1,507	12			1,507	12					21.1	(14)		
		1,486	20			1,483	21					21.7	(16)

MLE 4.8 (contd.)

	The second second second		The second secon							TO TOUT	, T		
s-cast	٦	700°C,	700°C, 30 d	As-cast	, t	700°C, 30 d	30 d				)		
	°1 %	d o	% I %	o o	% Iodo	o O	% I	5:19	6	1:5	2	2:17	7
469	20	1,469	27	1.473	20	20 1,473	26			20.2 (21)	(21)	40.5 32.4	(25)
.345	1	1,345	۵	1,350	15	1,363	on '	30,12	( 26)	30,1	(24)	22.6	
3.5		1,324	13			1,330	15	·				33,3	(15)
2	ì ; ;	1,248	16	37-10		1.248	11					41.6	(6)
243	10	1,239	1	1.241	12			22.1	(23)	22.0 (31)	(31)		

corresponding to the RE-rich phase appeared at a lower temperature for both the annealed alloy and the as-cast alloy. Though this phase could be accounted for 5:19 in the X-ray (Table 4.8) its Curie temperature is much lower (150°C) than that expected for a 5:19 phase (280°C). Also the MM-Co-Fe phase diagram (Figure 4.1) shows that the 5:19 phase does not extend beyond about 10 a/o Fe. It is possible that this phase is of 2:7 type rather than 5:19.

### 4.4 Conclusions

Phase equilibria in MM-Co-Fe system was studied; at 900°C between 73 a/o and 92 a/o Co with Fe content upto 15 a/o. Six different phases MMCo3, MM2Co7, MM5Co19, MMCo5, MM2Co17 and β-Co were identified close to the MM-Co binary line. Thermomagnetic analysis proved to be a better technique than X-ray diffraction for identifying the low cobalt phases, MMCo2, MM2Co7 and MM5Co19 the ferromagnetic transition temperature of which were found to be 20°C, 67°C and 270°C respectively. The 2:7, 5:19, 1:5 and 2:17 phases were also found to exist as stable phases in the MM-Co-Fe ternary system. The 5:19 phase was found to extend to about 10 a/o Fe while the 2:7, 1:5 and 2:17 phases extend to greater than 10 a/o Fe. In addition to the above phases, a phase with a T of 340°C was found to coexist with 5:19 and 1:5 phases close to the 5:19 stoichiometric line and the region of its occurrence in the MM-Co-Fe system at 900°C is indicated. The phase on the

cobalt rich side of 2:17 stoichiometry had  $\beta$ -Co structure with no solid solubility for RE elements while MMCo<sub>5</sub> showed a narrow homogeneity region at 900°C other phases did not show the presence of such region. MMCo<sub>5</sub> was unstable at temperatures below about 700°C. The decomposition was found to be sluggish in the low Fe (<3 a/o) containing 1:5 phase and hence evidence could be obtained only for the MM<sub>2</sub>Co<sub>17</sub> product phase. But in high iron 1:5 alloys (>5 a/o) the decomposition at 700°C was rapid and evidence could be obtained for both the 5:19 and 2:17 product phases by TMA.

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### 5. MAGNETIC PROPERTIES OF MM\_Co\_Fe PHASES

### 5.1 Introduction

Primary magnetic properties of interest for permanent magnet production are (1) saturation magnetisation ( $4\pi\,\mathrm{M}_{\mathrm{S}}$ ) (2) magnetocrystalline anisotropy field ( $\mathrm{H}_{\mathrm{A}}$ ) and (3) Curie temperature. Materials with high values for all these parameters are excellent candidates for the production of high energy density permanent magnets. Hence it is very essential to characterise the different intermetallic phases occurring in the RE-Co systems for these parameters. Extensive studies have been made on the primary magnetic properties of pure binary RE-Co phases and the properties reported for the intermetallic phases in the RE-Co systems RE-Ce, La, Nd, Pr and Sm are summarised in Tables 6.1 and 6.2.

### 5.2 Experimental

The different intermetallic phases found in the MM-Co-Fe system were characterised for their saturation magnetisation and Curie temperature. The experimental procedure is the same as that described under Section 3.2.3. The H<sub>A</sub>, as explained under the Section 3.3.1.1 was not determined in this study.

TABLE 5.1. PRIMARY MAGNETIC PROPERTIES OF RECo5 PHASES

System		1	1	1	I v	† 2 1	
Properties	Unit	YCo <sub>5</sub>	LaCo <sub>5</sub>	CeCo <sub>5</sub>	CeMMCo	PrCo <sub>5</sub>	SmCo <sub>5</sub>
4π M <sub>S</sub> (20°C)	G	10600	9090	7700	8900	12000	9650
T <sub>C</sub>	°C	<b>64</b> 8	567	374	495	612	724
H <sub>A</sub> (20°C)	KOe	130	175	170-210	180-195	145-210	210-290
(BH) <sub>max</sub> (Theoretical)	MGOe	28.1	20.6	14.8	19.8	36.0	23.4
i <sup>H</sup> c	0e	6700	3600	2800	4750	5 <b>7</b> 50	25000

TABLE 5.2. MAGNETIC SATURATION AND ANISOTROPY AT 20°C OF RE<sub>2</sub>Co<sub>7</sub> PHASES

Phase	4π M <sub>s</sub>	(20°C)	H <sub>A</sub> (KOe)
	(emu/g)	Gauss	A
La <sub>2</sub> Co <sub>7</sub>	50.4	5193	171
Pr <sub>2</sub> Co <sub>7</sub>	83.4	8851	105
Nd <sub>2</sub> Co <sub>7</sub>	85.1	9173	28

## 5.3. Results and Discussion

# 5.3.1 Saturation Magnetisation of MM-Co Phases

The specific saturation magnetisation ( $\sigma$ , emu/g) was determined for the stoichiometric phases in the MM-Co system. The amount of the second phase in these alloys was 5% as determined metallographically. The  $4\pi$  M of MM<sub>2</sub>Co<sub>7</sub>, MM<sub>5</sub>Co<sub>19</sub> and MM<sub>2</sub>Co<sub>17</sub> phases are shown in Table 5.3.

TABLE 5.3. SATURATION MAGNETISATION OF MM\_Co PHASES AT 20°C

Alloy	Phase		omposition o, analyse		4π M	4π M	Mr/Ms
		RE	Со	Fe	(emu/g)	(emu/g)	
A-3	2:7	22.9	73.4	3.7	29	12	0.41
B-3	5:19	21.5	75.0	3.5	44	29	0.66
C-3	1:5	16.9	80.3	2.8	95	80	0.84
D-4	2:17	11.1	87.1	1.8	114	21	0.18

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The MMCo $_5$  phase has a  $4\pi\,\mathrm{M}_\mathrm{S}$  of 95 emu/g and a density of 8.3 g/cc. This on conversion turns out to be 9.6 KG which is slightly higher than the value reported by  $\mathrm{Strmat}^1$  for  $\mathrm{CeMMCo}_5$  phase (Table 5.1). The  $4\pi\,\mathrm{M}_\mathrm{S}$  of the individual RECo $_5$  phases differ considerably (Table 5.1). The higher value observed for the  $\mathrm{MMCo}_5$  may be due to a different distribution of RE elements in the Indian mischmetal and due to the presence of about 6 w/o Fe as impurity in the Indian mischmetal.

The MM<sub>5</sub>Co<sub>19</sub> has a  $4\pi$  M<sub>s</sub> of 44 emu/g. Though it contains 75.0 a/o Co its  $4\pi$  M<sub>s</sub> is low by more than twice that of MMCo<sub>5</sub> which contains 80.3 a/o Co. The MM<sub>2</sub>Co<sub>7</sub> phase which contains 73.4 a/o Co shows a  $4\pi$  M<sub>s</sub> of 29 emu/g only. However no  $4\pi$  M<sub>s</sub> value is available for Ce<sub>5</sub>Co<sub>19</sub> and Ce<sub>2</sub>Co<sub>7</sub> to compare with the observed values. Both Ce<sub>5</sub>Co<sub>19</sub> and Ce<sub>2</sub>Co<sub>7</sub> are paramagnetic at room temperature (25°C). The T<sub>c</sub> of Ce<sub>5</sub>Co<sub>19</sub> is 20°C and that of Ce<sub>2</sub>Co<sub>7</sub> is -110°C.<sup>4</sup>

The  $\text{MM}_2\text{Co}_{17}$  phase contains 87.1 a/o Co and shows the highest value for  $4\,\pi\,\text{M}_{_{\rm S}}$  as expected but on the contrary it has the lowest  $\text{M}_{_{\rm T}}/\text{M}_{_{\rm S}}$  ratio of 0.18.  $\text{MM}_2\text{Co}_{17}$  phase has been reported to have an easy plane anisotropy 5 in which case this low value is explainable.

## 5.3.2 Curie Temperature

5.3.2.1 T of Stoichiometric Phases in MM-Co System

The chemical and phase composition of the alloys used for the thermomagnetic study are given in Table 5.4.

TABLE 5.4. CHEMICAL AND PHASE COMPOSITION OF MM\_Co-Fe ALLOYS USED FOR TMA

			•		
Alloy	Composi	tion (a/o,	analysed)	Phases (900°	c, 4-10 d)
	RE	Co	Fe	Microstructure (number of phases)	I I
A-1	23.4	72.8	3.8	2	1:3 + 2:7
6-1	22.3	72,3	5.4	2	2:7 + 5:19
12-1	21.3	67.3	11.4	2	2:7 + 5:19
15-1	21.5	63,3	15.2	2	2:7 + 5:19
B-5	21.0	75.6	3.4	2	5:19 + 1:5
C-1	19.5	77.4	3.1	2	5:19 + 1:5
C-3	18.1	79.0	2.9	2	5:19 + 1:5
S-1	16.5	83.5	0	1	1:5
0-3	16.6	83.4	0	1	1:5
6-5	16.6	77.0	6.4	1	1:5
12-5	15.8	71.9	12.3	2	1:5 + 2:17
			1 - 20		

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The thermomagnetic curves of alloys A-1, B-5 and S-1 are shown in Figure 5.1. A magnetic transition near 60°C is observed for alloy A-1. Since the composition of this alloy lies on the MM rich side of 2:7 stoichiometry this transition is attributed to the  ${\rm MM}_2{\rm Co}_7$  phase. The second phase seen in the microstructure of the alloy should be the 1:3; hase. As this phase is expected to have a  $T_{C}$  much less than room temperature (20°C) its transition could not be observed in the present study. The alloy B-5 with composition on the cobalt rich side of 5:19 stoichiometry shows no magnetic transition near 60°C confirming the absence of a 2:7 phase. A major fall in moment observed at 280°C is attributed to the 5:19 phase. For the alloy S-1, a magnetic transition corresponding to the 1:5 phase only is observed. The T values of these phases after correcting for the thermal hysteresis are compared in Table 5.5. with the literature values for low cobalt phases in binary RE-Co systems. It is interesting to note that a simple relation i.e.

$$T_{c \text{ MM}_{\mathbf{x}}Co_{\mathbf{y}}} = [ (wt \% \text{ RE in MM})(T_{c \text{ RE}_{\mathbf{x}}Cc_{\mathbf{y}}})] 10^{-2}$$

could be used to predict the  $T_{C}$  of a MM  $Co_{Y}$  phase from the composition of MM and the  $T_{C}$  of binary  $RE_{X}Co_{Y}$  phases. For example, this equation when applied to the alloy S-1 (Table 5.5) which was prepared with high purity RE elements and  $Co_{Y}$ 

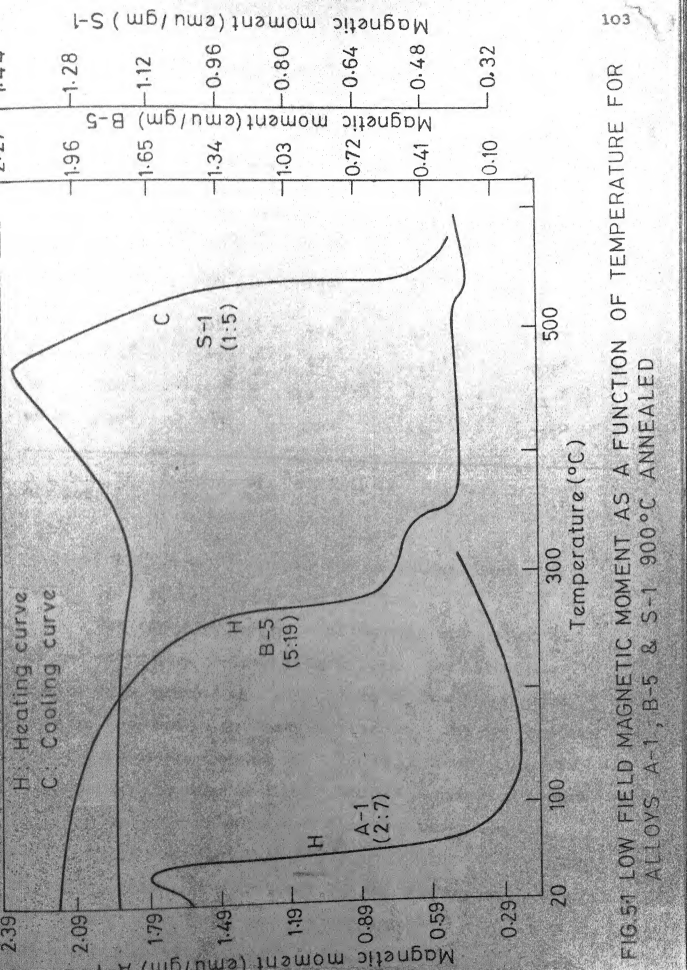


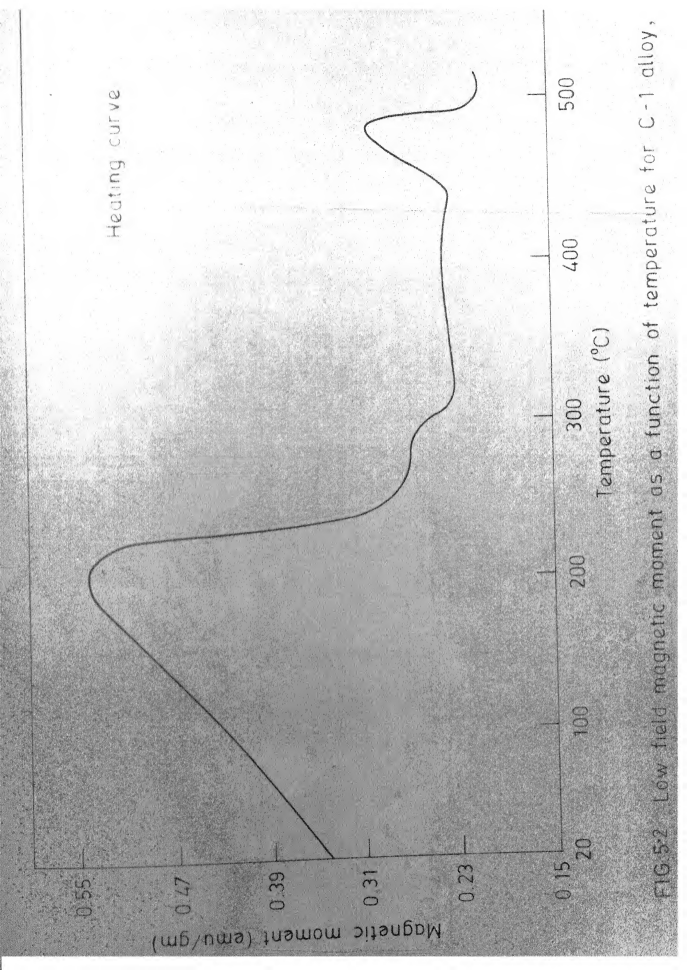
TABLE 5.5. CURIE TEMPERATURES IN °C OF RE-CO COMPOUNDS

RE	RECO <sub>3</sub>	RE <sub>2</sub> Co <sub>7</sub>	RE <sub>5</sub> Co <sub>19</sub>	RECO <sub>5</sub>	RE <sub>2</sub> Co <sub>17</sub>
Ce	<b>-</b> 195 <sup>a</sup>	-110 <sup>C</sup>	20 <sup>d</sup>	374 <sup>e</sup>	810 <sup>£</sup>
La	Does not exist	220 <sup>C</sup>	343 <sup>d</sup>	567 <sup>e</sup>	Does not exist
Nd	122 <sup>a</sup>	355 <sup>C</sup>	441 <sup>d</sup>	630 <sup>C</sup>	877 <sup>£</sup>
Pr	76 <sup>a</sup>	337 <sup>C</sup>	417 <sup>d</sup>	612 <sup>e</sup>	898 <sup>£</sup>
Sm	250 <sup>b</sup>	450 <sup>b</sup>	(510?) <sup>b,d</sup>	724 <sup>e</sup>	917 <sup>£</sup>
MM	< 20 <sup>p</sup>	67 <b>P</b>	270 <sup>p</sup>	541 <sup>p</sup>	>735 <sup>p</sup>

a: Ref. 7; b: Ref. 8; c: Ref. 4; d: Ref. 3; e: Ref. 1; f: Ref. 9; p: present work.

predicts a T<sub>c</sub> of 491°C, whereas the experimental value is 541°C, the difference being less than 10%.

For the alloy B-5 two more magnetic transitions at 350 and 517°C were observed (Figure 5.1). The latter was due to MMCos present as a minor phase in this alloy which was also detected in the X-ray diffraction. The new magnetic transition observed between the T of MM5Co19 and MMCo5 was also seen for a number of alloys near 1:4 composition in the MM-Co system 10 and this has been discussed under Section 4.3.1. It is seen from Figures 5.2 and 5.3 that the heating and cooling curves for the alloy C-1 are irreversible. In the heating cycle the magnetic moment increases steadily and peaks near the Curie temperature before falling. On cooling



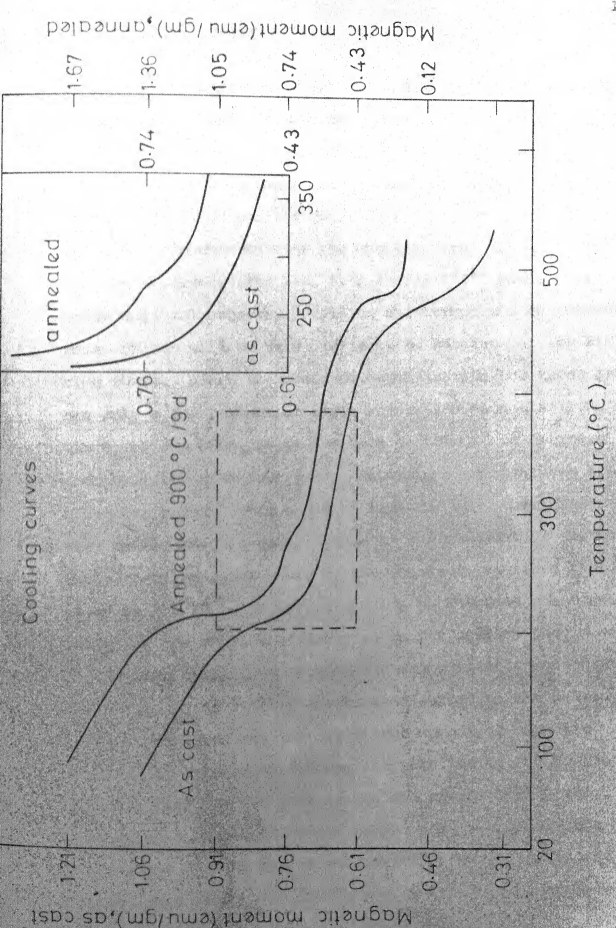


FIG.53 LOW FIELD MAGNETIC MOMENT AS A FUNCTION OF TEMPERATURE FOR C-1 ALLOY

it takes a different path and the magnetisation at any given temperature is greater than that observed in the heating This behaviour was observed for most of the alloys cycle. studied and in particular for the as-cast alloys. In subsequent heating and cooling cycles the curves were reversible and the peeking phenomena near the magnetic transition temperature was not observed. The 2:7, 5:19 and 1:511,12 phases have large uniaxial magnetocrystalline anisotropy and consequently these phases will be very difficult to saturate at low fields like 40 Oe. Hence the observed magnetisation for these samples are only a few percent of their saturation magnetisation. The  $H_A$  of the MMCo<sub>5</sub> phase has been reported 13 to decrease monotonically with increasing temperature and vanishes near its Curie point. This might be true for the other phases also in the MM-Co system. Hence as the temperature approaches the transition point where HA also tends to vanish, a field of 40 Oe should be sufficiently high to increase the magnetisation of the samples giving rise to the peaking phenomena in the heating cycle. On cooling the magnetisation was observed to increase 2 to 3 times the value observed in the heating cycle. This suggests that increased number of magnetic domains get aligned in a direction parallel to the applied magnetic field in the polycrystalline sample during the cooling as a result of domain growth. In the subsequent heating and cooling cycles the curves were reversible and the peaking phenomenon near the Tc was absent. This suggests

that the magnetic domain configuration remains essentially the same after the first heating and cooling cycles.

The Curie temperature of the  ${\rm MM}_2{\rm Co}_{17}$  phase is not reported here as it has a T $_{\rm C}$  >700°C, which is beyond the experimental safety limit of the high temperature oven used.

## 5.3.2.2 T<sub>c</sub> of MM-Co-Fe Phases

The compositions of the alloys A-1,35,6-1, 12-1 and 15-1 lie between the 2:7 and 5:19 stoichiometries. microstructure of the alloys show 2 phases (Table 5.4). These phases are assigned to MM2Co7 and MM5Co19. The heating TMA curves of the alloys A-1, 6-1, 12-1 and 15-1 are shown in Figure 5.4. Their Curie temperature after correcting for the thermal hysteresis are indicated in Table 5.6. It could be seen that the Curie temperature of both the 2:7 and 5:19 phases increase with increasing concentration of Fe. A three fold increase in the Tc of the 2:7 phase, from 70°C to 218°C, is observed as the Fe content varies from 3.8 a/o in alloy A-1 to 15.2 a/o in alloy 15-1 (Table 5.6). On the contrary only about 60°C raise in T is observed for the 5:19 phase as the Fe varies from 3.8 to 11.4 a/o (Figure 5.5). It is to be noted that the concentration of Fe listed in Table 5.6 is only the bulk concentration of the alloy and the actual distribution of the Fe in the two phases when they coexist, need not be the same. Thus the observed lower raise in T for the MM5Co19 could possibly be due to the lesser solubility of Fe in it than in the 2:7 phase. There is also indirect indication

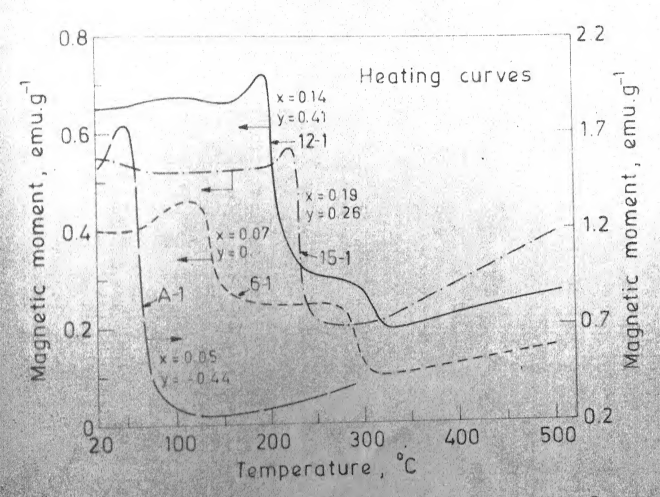


Fig 54 Magnetisation versus temperature at 40 Oe for MM<sub>2</sub>(Co<sub>1-x</sub>Fe<sub>x</sub>)<sub>7+y</sub> alloys; 900 °C, 7 days annealed

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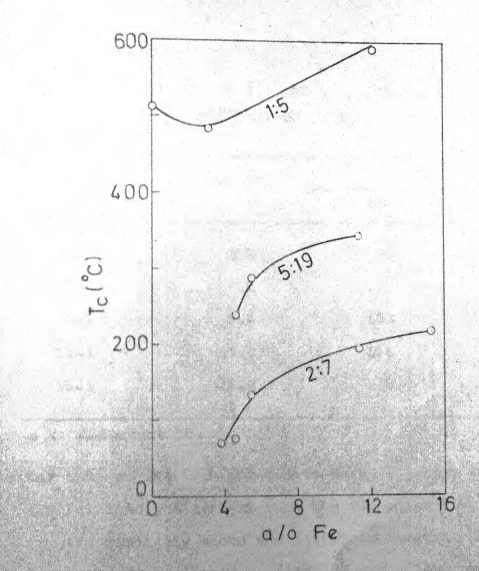


FIG.55 VARIATION OF To OF 2:7 5:19 AND 1:5:

PHASE WITH FE CONCENTRATION IN

MM-Co-FE SYSTEM

TABLE 5.6. T<sub>C</sub> OF THE 2:7 AND 5:19 PHASES WITH THE VARYING FE CONCENTRATION IN THE MM-CO-FE SYSTEM

Alloy	a/o Fe	Tc	(°C)
		2:7	5:19
A-1	3.8	<b>7</b> 0	absent
36 <sup>a</sup>	4.6	<b>7</b> 5 .	238
6-1	5.4	133	280
12-1	11.4	196	295
15-1	15.2	218	absent

a: Reference 10.

for this as the 5:19 phase extends to a lesser extent (~11.4 a/o Fe) into the ternary than the 2:7 phase (Section 4.3.1).

Similarly along the 1:5 stoichiometric line alloys 0-3, 6-5 and 12-5 were studied. These alloys contain different amount of Fe and their microstructure revealed (Table 5.4) that the alloys 0-3 and 6-5 to be single phase 1:5 while 12-5 contains some amount of other phase ( 20%) in addition to the major 1:5 phase. The TMA plots of these alloys are shown in the Figure 5.6 and their T<sub>c</sub>scorrected for the thermal hysteresis are shown in Table 5.7. The T<sub>c</sub> initially falls to 485°C at 2.9 a/o Fe from 512°C and then rises to 580°C at 12.3 a/o Fe (Figure 5.5).

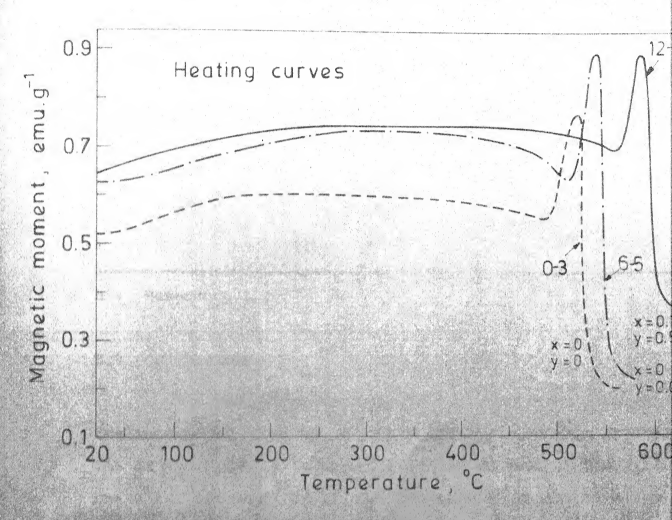


Fig.5:6 Magnetisation versus temperature at 40 De for MM(Co<sub>1 x</sub>Fe<sub>x</sub>)<sub>5.y</sub> alloys

TABLE 5.7. T OF THE 1:5 PHASE WITH THE VARYING FE CONCENT-RATION IN THE MM-Co-Fe SYSTEM

Alloy	a∕o Fe	T <sub>C</sub> (°C)
0-3	0	512
C-3 <sup>a</sup>	2.9	485
<b>6- 5</b>	5.4	532
12-5	12.3	580

a: Reference 11.

### 5.4 Conclusions

The saturation magnetisation and Curie temperature of the intermetallic phases in the MM-Co-Fe system were studied. The  $4\pi$  M<sub>S</sub> of the 1:5 phase is 95 emu/g and that of the 2:17 phase is 114 emu/g. Both the 2:7 and 5:19 phases have a value less than 45 emu/g. The T<sub>C</sub> of the stoichiometric 2:7, 5:19 and 1:5 phases are 67°C, 270°C and 541°C respectively. These values were found to increase with the increasing concentration of Fe.

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OF MM(Co<sub>1-x</sub>Cu<sub>x</sub>)<sub>5</sub> ALLOYS

### 6.1 Introduction

Widespread application of  $SmCo_5$  permanent magnets is limited by the high cost of Sm and the scarcity of cobalt. Attractive magnetic properties have been reported for  $SmCo_5$  alloys containing  $Cu.^{1-2}$   $MM-Co^3$  and  $MM-Co-Fe-Cu^4$  alloys also have useful permanent magnet properties of technological importance, but no systematic studies on phase relations and magnetic properties of MM-Co alloys containing Cu have been reported. Hence a systematic addition of Cu in place of Co was made in  $MMCo_5$  alloy to know (i) the extent of solid solubility of Cu in  $MMCo_5$  (ii) its effect on the phase stability of the 1:5 phase and (iii) its effect on  $4\pi M_S$  and  $T_C$  of 1:5 phase. The mischmetal used in this study has the same composition as given under Section 3.1.1.

## 6.2 Experimental

The melting and annealing of the alloys and the procedure of phase analysis are the same as described under Sections 3.1 and 3.2. In X-ray diffraction studies, for Cu rich alloys Cu  $K_{\rm CC}$  radiations gave better results than Co  $K_{\rm CC}$  radiation. The magnetisation and the Curie temperature were studied with a vibrating sample magnetometer. To measure the  $4\pi$  M<sub>s</sub>, the sample were powdered, magnetically aligned

and then fixed in an epoxy resin. A field of 11 KOe was sufficient to saturate the aligned samples. The magnetisation of the alloys as a function of temperature was studied in a low magnetic field of about 40 Oe in order to determine their magnetic phase transitions. Solid pieces of about 0.1 gm were used for this study. The experimental set up is shown in Figure 3.2 and the procedures/given under Section 3.3.1.3.

### 6.3 Results and Discussion

### 6.3.1 Phase Relations

The CaCu<sub>5</sub> type phase is present in the entire composition range in MM(Co<sub>1-x</sub>Cu<sub>x</sub>)<sub>5</sub> alloys (x = 0 to 1) (Table 6.1). The low Cu alloys B-1 and B-2, annealed at 900°C contain 2:17 and a new phase in small amounts in addition to the major 1:5 phase. The amount of 2:17 phase increases when these alloys are annealed at  $700^{\circ}$ C. The X-ray diffraction patterns of alloys B-4 and B-5 show 1:5 phase both in  $900^{\circ}$ C and  $700^{\circ}$ C annealed condition whereas their microstructure gives a clear evidence for the presence of a small amount of a second phase also, which gets etched brighter than the matrix. Perry<sup>5</sup> reports that in Sm-Co-Cu system SmCo<sub>5</sub> phase coexists with a Co phase on the transition metal rich side of 1:5 stoichiometry for alloys containing greater than 10 a/o Cu at 800°C. It is possible that this phase is Co.

CHEMICAL AND PHASE COMPOSITION OF MM1-z(Co1-x-y Fe Cux) ALLOYS

Alloy		MM1-Z (Co1-x-y	y Fe	Com	Composition (w/o,	w) uo	10,		Phases (	Phases (X-ray, microstructure)	ostn	cture)
	gx/2			ana	analysed)				900°C, 4 d	þ	7	700°C, 4 d
	×	<b>&gt;</b>	<b>N</b>	RE	ပ္ပ	다 0	ਰ	Mic. St.	X-ray	TMA (T°C)	Mic. st.	X-ray
ä	; •	0.02	0.07	29.3	60.7 2.2	2.2	7.8	е	1:5+2:17+x	550, ~100	٣	1:5+2:17+x
B-2	0.2	0.02	0.07	29.2	53.8	2.1	14.9	ю	1:5+2:17+x	500, 100	ო	1:5+2:17+x
B-3	e 0	0.03	°°o	31.6	44.7	2.2	21,5	- 0	1:5+x	320,(~100)		Not observed
B-4	0.4	0.02	0.07	29.5	38.4	2.1	30.0	7	1:5+(Co)	230	7	1:5+(Co)
B-5	9.0	0.02	0.07	29.5	25.1	2.1	43.3	7	1:5+(Co)	₹20	7	1:5+(Co)
B-6	0	0.02	0.07	28.7	12.2	2.1	57.0	т	1:5+1:6+(Co) <20	₹20	7	1:5+(Co)
B-7	1.0	0,02	0.07	28.3	0	1.9	8.69	2	1:5+1:6		7	1:5+1:6
							- XXII	- *				

However, it could not be confirmed by X-ray since it is present in small amounts. The alloys B-6 and B-7 melted on heating at 900°C and when quenched from 900°C showed 1:5 and 1:6 phases. This is similar to a CeCus alloy which at 900°C has a liquid phase in equilibrium with a solid phase (1:6). The MMCu<sub>6</sub> phase has a SmCu<sub>6</sub> type structure. The orthorhombic lattice constants of MMCu, are a: 8.132 %,  $\underline{b}$ : 5.099  $\mathring{A}$  and  $\underline{c}$ : 10.211  $\mathring{A}$ . On annealing at 700°C the 1:6 phase observed in alloy B-6 disappears giving rise to 1:5 phase, and a trace of a third phase (Co) observed at 900°C remains at 700°C. The alloy B-7 which contains no Co still shows 1:5 and 1:6 phases at 700°C. In binary Ce-Cu<sup>6</sup> system these phases exist as line compounds with no homogeneity region and a slight deviation from the stoichiometry will result in two phase structure. The MMCu, and MMCu, phases differ by less than 4 w/o in their stoichiometries and the presence of two phase structure observed for the B-6 alloy is attributed to its slightly transition metal richer composition than 1:5. The lattice constants a and c of the 1:5 phase in alloys B-1 to B-7 increase linearly with increasing Cu concentration (Figure 6.1). This is similar to the results reported by Lihl<sup>8</sup> for YCo5-YCu5 and SmCo5-SmCu5 systems.

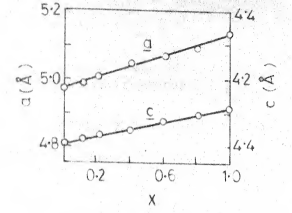


FIG. 6-1 LATTICE PARAMETERS VS COMPOSITION
OF 1:5 PHASE

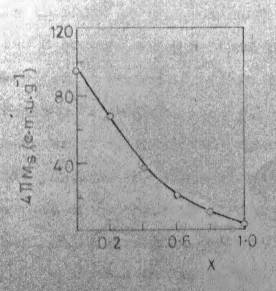
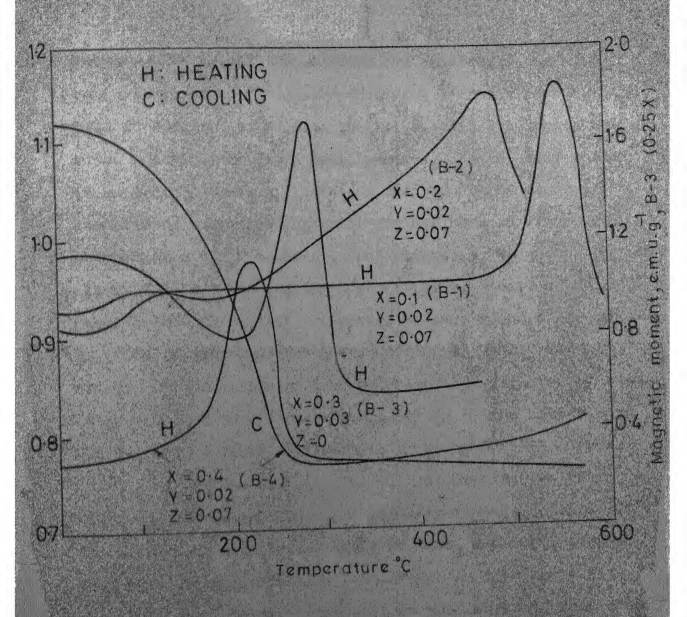


FIG FZ 4TMs VS COMPOSITION OF

# 6.3.2 Magnetic Characteristics

The replacement of Co by Cu in MMCo $_5$  lowers the  $4\,\pi\,{\rm M}_{\rm S}$  from 95 emu/g to less than 20 emu/g at 50 w/o Cu (Figure 6.2).

Alloy B-1 which contains 7.5 w/o Cu shows a Curie temperature greater than 500°C (Figure 6.3). MM-Co-Cu alloys close to this composition have been studied by Walkiewicz et.al.4 and they reported an addition of 0.2 w/o Mg to alloys near this composition increases the  ${}_{i}{}^{H}{}_{C}$  to greater than 24 KOe. On further increasing the Cu content the Tc falls to 500°C at 14.5 w/o Cu (Figure 6.3) for the alloy B-2, to 320°C for the alloy B-3 and to 230°C for the alloy B-4 (Table 6.1). Beyond 15 W/o Cu the fall in  $\text{T}_{\text{C}}$  is steeper with increasing Cu content. For all the alloys two common features were observed in their TMA curves measured in a low magnetic field: firstly the shape of the cooling curve was different from that of the heating curve, secondly the magnetisation of the sample increased manyfold on cooling. This is shown for the alloy B-4 in Figure 6.3. This was observed for the Cu free MM-Co phases also and it was attributed to the strong temperature dependence of HA of the phases in those alloys.9 The Ha of the 1:5 phase decreases with increasing temperature and vanishes near the Curie point. 10 Hence at a fixed magnetic field the magnetisation should be easier at temperature close to the Curie point. A temperature difference of about 10°C is present between the heating and cooling curves



FOR MM<sub>1-z</sub> (Co<sub>1-x-y</sub> Fey Cux ) ALLOYS, 900 °C

because of a thermal lag between the sample and the high temperature oven. The microstructures of the alloys B-1 and B-2, in addition to showing the 1:5 contain 2:17 and a new phase (darkly etched) in small amounts. As the 2:17 phase is expected to have a T<sub>C</sub> greater than 600°C the magnetic transitions corresponding to it are not seen in Figure 6.3; whereas near 100°C there are indications in TMA curves of the alloys B-1, B-2 and B-3 for a third phase. Based on X-ray results Labulle et.al. 11 reported to have found a new phase Ce<sub>5</sub>(CoCu)<sub>13</sub> and a hexagonal Ce<sub>5</sub>(CoCu)<sub>19</sub> in CeCo<sub>3.5</sub>Cu<sub>1.5</sub> alloy. However, no magnetic data have been reported by them. It is possible that the new phase may be of the latter type but it needs further confirmation by X-ray and EPMA study.

Alloys B-5 and B-6 were also studied by TMA upto 600°C and they do not show any magnetic transition above 20°C. The alloy B-7 which is a mixture of MMCu<sub>5</sub> and MMCu<sub>6</sub> phases show non-zero magnetisation (~3 emu/g) at 20°C while CeCu<sub>5</sub><sup>12</sup> and CeCu<sub>6</sub><sup>13</sup> are paramagnetic at room temperature. This is attributed to Fe present as an impurity in the mischmetal used in this work.

#### 6.4 Conclusion

The substitution of Cu for Co in MM( ${\rm Co}_{1-x}{\rm Cu}_x$ )5 alloys gives stable 1:5 phase in the entire range of composition. At low Cu concentration (21.5 w/o) 2:17 and a new

phase coexist with the matrix 1:5 phase. Cu could be substituted for Co upto 15 w/o in 1:5 with a T $_{\rm C}$  not less than 500°C and  $4\,\pi\,{\rm M}_{\rm S}$  of about 70 emu/g. Beyond 15 w/o Cu both  $4\,\pi\,{\rm M}_{\rm S}$  and T $_{\rm C}$  are considerably reduced.

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#### 7. PERMANENT MAGNET PROPERTIES OF 1:5 TYPE MM-Co ALLOYS

#### 7.1 Introduction

Previous work on the development of mischmetal cobalt permanent magnets and the aims of the present work are given below.

## 7.1.1 Previous Work on MMCo5 Magnets

The first successful results on the permanent magnet properties of sintered MM-Co magnets were reported by Fellows et.al. 1 Subsequently Nagel 2-3 and Narita 4-5 have also reported the properties achieved for MMCo5 magnets. The materials used, the processes followed and the properties developed for MMCo, by them are summarised in Table 7.1. It could be seen from the table that the composition of MM used and the processes adopted for the fabrication of magnets are different in each case. Also the alloy composition used by Narita and Fellows et.al. is close to MM<sub>5</sub>Co<sub>19</sub> stoichiometry which on sintering results in a predominantly MMCo5 phase magnet. The former used liquid phase sintering whereas the latter followed solid phases sintering on alloys of single composition without low melting sintering additive. The Br and (BH) wax values were not reported by Narita. He proposes that the large scale shift in composition should have occurred during sintering due to the loss of RE.

*				
Material	Composition	Process	Properties	Reference
Cerium mischmetal and Co (RE dist- ribution in MM and the purity of Co are not given)	64.5 w/o Co, 35.5 w/o MM (base); 27.0 w/o Co, 73.0 w/o MM (sinter additive). Aimed composition of the mixture of the powders 1 and 2 is 59.8 w/o Co and 40.2 w/o MM	Arc melted, attrition milled in petroleum ether, aligned in a pulse field of 50 KOe, axially pressed to 21 kbar in a holding field of 7 KOe and finally isostatically pressed to 7 kbar in the absence of field, sintered between 990 and 1110°C for	P: 8.0-8.2 g/cc i <sup>H</sup> : 4-6 KOe B : 4-7 KG (BH) max MGOe	Fellows et al., 1972 (1)
Natural mischmetal (53 w/o Ce, 30 w/o La, 13 w/o Er) and 4 w/o Er) and cobalt (99.9% pure)	Alloys on RE rich side of 1:5 stoi- chlometry were used. Actual com- position studied is not given	Induction melted, jet milled with nitrogen, aligned in 50 KOe field, isostatically pressed at 6000 atm. Sintered at 1040°C and heat treated. Oxide content of the sintered magnet was ~ 0.5 w/o	the theoretical density (8.3 g/cc) i H : 9 KOe Br : 8.1 KG (BH) max MGO	Nagel et al., 1975, 1976 (2-3)
Mischmetal (49.5 MMCo <sub>x</sub> alloys % Ce, 20.2% La, 366 x & 4.8 18.8% Nd, 5.0% Pr, Were used 1.9% Sm, 3.9% Fe, 0.03% Mg and the balance impurities unidentified) and cobalt (99.5% Co, 0.15% Ni and 0.14% Fe), all in w/o	MMCo <sub>x</sub> alloys  3. Were used 4.8  3. W/o	Arc melted, ball milled in acetone, aligned in 10 KOe, pressed further in the absence of field, sintered between 960 and 1030°C with no sinter additives	1, c. 8, 3 Koe	Narita et al.

## 7.1.2 Aims of the Present Work

Using commercial grade Indian mischmetal and cobalt (compositions are same as given under Section 3.1.1) work was undertaken (i) to develop the process: comminution, compaction and sintering procedures to fabricate MM=Co magnets (ii) to correlate the properties of the sintered magnets to their structure and (iii) to identify the problem - areas in the development of MM=Co permanent magnets.

#### 7.2 Experimental

The experimental procedure for the magnet processing are described in detail in Chapter 3 under Section 3.3 and the steps involved are illustrated in Figure 3.3 together with the materials and the equipments used. The compositions of the alloys studied are given in Table 7.2.

### 7.3 Results in the course of t

The permanent magnet properties of MM-Co alloys were studied under three different conditions: (1) Resin bonded sample, (2) powder compacts and (3) sintered pellets. In the case of sintered pellets, the properties obtained were monitored by studying their structure by X-ray diffraction.

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# 7.3.1 Resin Bonded MM-Co Magnets

Prior to compaction and sintering it was necessary to select an appropriate composition and milling time that

TABLE 7.2 COMPOSITION OF MM-Co ALLOYS USED FOR MAGNET FABRICATION

Alloy formula	Composi	tion (w/o, nom	inal)
	MM	Co	Fe*
MMCo3.5	40.9	56.5	2.6
MMCo3.8	38.8	58.7	2.5
MMCo4.0	37.5	60.1	2.4
MMC04.2	36,5	61.2	2.3
MMC04.4	35.4	62.4	2.2
MMC04.6	34.4	63.4	2.2
MMCo <sub>4.8</sub>	33.3	64.6	2.1
MMCo5.0	32.5	65.4	2.1
MM* Co3.8	31.4 MM 8.1 Sm	58,5	2.0
MM* CO4.0	30.3 MM 7.7 Sm	60.1	1.9

Fe . The impurity Fe in natural mischmetal is substracted (6 w/o assumed) and is shown separately.

give powders with maximum coercivity. For this purpose  ${\rm MMCo}_{\rm X}$  alloys with X lying between  ${\rm MMCo}_{3.5}$  and  ${\rm MMCo}_{5}$  were milled in 1 gm quantity in a plastic container upto 20 hours. The powder was mixed with an epoxy resin in a definite ratio, aligned in a magnetic field and set in air. The  ${}_{\rm i}{}^{\rm H}{}_{\rm C}$  and  $4\pi$  M of these resin bonded magnets were measured and the results are shown in Figure 7.1. A maximum  ${}_{\rm i}{}^{\rm H}{}_{\rm C}$  of 3100 Oe

MM\*, The mischmetal contains 20 w/o Sm added intentionally.

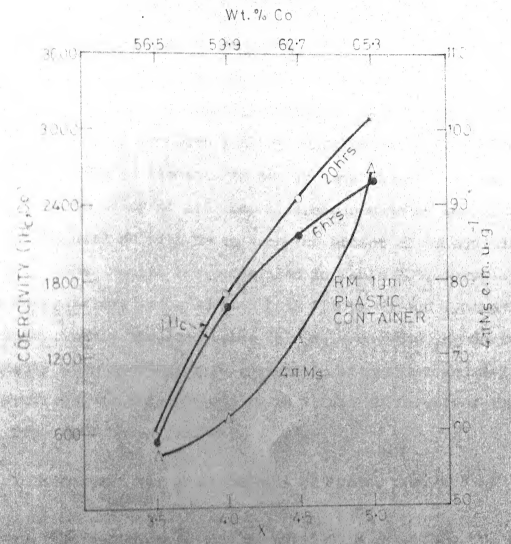


Fig. 74  $\pm$  mMs and iHs for MMCoX althous  $3.5 \le X \le 5.0$ 

could be obtained for 20 hours milled MMCo $_5$  alloy and it has a  $4\pi$  M $_s$  of 95 emu/g. Alloys on MM rich side of MMCo $_5$  all have lower values for  $_{1}^{H}_{C}$  and  $4\pi$  M $_s$ .

In order to know the effect of Fe and other impurities present in natural mischmetal (Flints MM) on the  $_{i}^{H}{}_{C}$ , MMCo $_{5}$  alloys were prepared with synthetic mischmetal containing pure ( $\geqslant$ 99.9%) RE elements in nearly same ratio as in natural MM. Also an alloy of 1:5 composition was made by substituting 20% of natural MM with Sm to know the effect of Sm addition on  $_{i}^{H}{}_{C}$ . The results are summarised in Table 7.3. Surprisingly the MMCo $_{5}$  alloys prepared from both Flints MM and Synthetic MM show exactly the same value for  $_{i}^{H}{}_{C}$ . But the  $_{i}^{H}{}_{C}$  decreases markedly with increasing Fe content in alloys S-1 and S-2. An arbitrary addition of 20% Sm to Flints MM increases the  $_{i}^{H}{}_{C}$  from 3200 oe to 4950 Oe.

## 7.3.2 Permanent Magnet Properties of Powder Compacts of MM-Co Alloys

The permanent magnet properties,  $M_r/M_s$ ,  $i^Hc'$ ,  $b^Hc$  and  $(BH)_{max}$  of field pressed powder compacts were measured for MM  $Co_x$ , 4.2 < x < 4.8, alloys in order to select the best composition giving maximum value for  $(BH)_{max}$ . Five grams of the alloys were milled for 3 to 9 hours as described under Section 3. to make 2 or 3 pellets each weighing about 2 gms. The properties measured for these green pellets are shown in Figure 7.2. The green density of the pellets were between 4.7 and 5.0 g/cc. The  $(BH)_{max}$  is highest for 6 hours milled

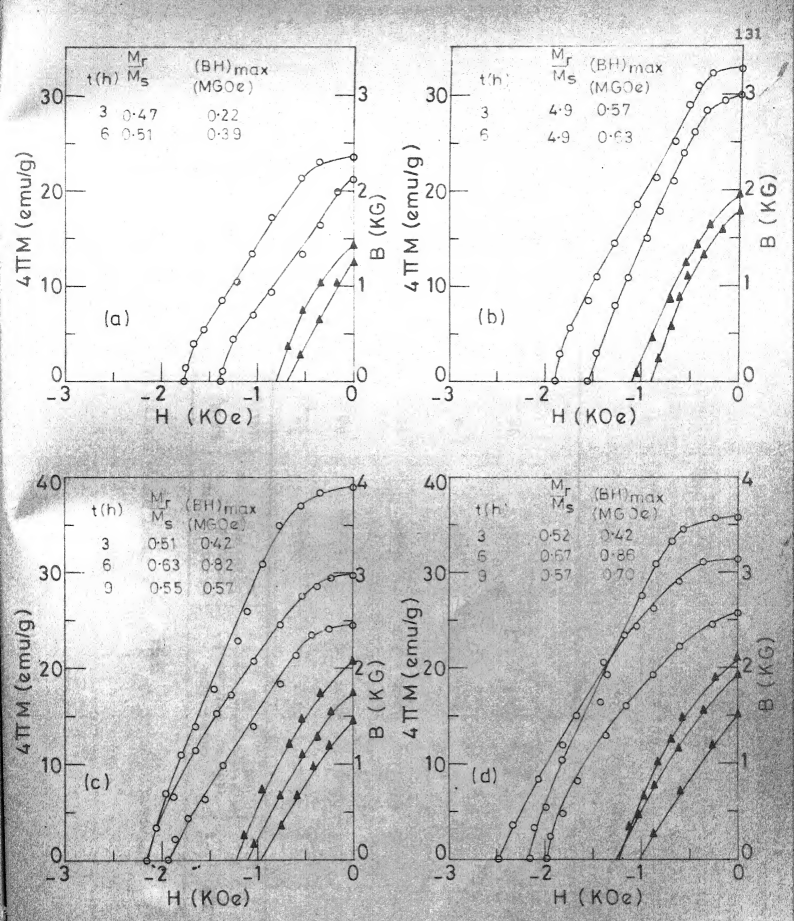


FIG.7-2 INTRINSIC AND INDUCTION DEMAGNETISATION CURVES FOR (a) MMCo42 (b) MMCo44 (c) MMCo44 AND (d) MMCo48

POWDER COMPACTS

COMPARATIVE STUDY OF 1 C OF DIFFERENT RESIN BONDED MMCOS MAGNETS TABLE 7,3

Alloy	Compc (w/o,	Composition /o, nominal)	n al)	Microstnucture (amount of	4π M S	Mr/Ms i H	r, H	Variable
4	MM	Co	Fe	%) yie piidse iii	(emu/g)		(oe)	
MMCo <sub>5</sub> (F)	32.8	65.1	2.1	2-10	0*96	08*0	3200	Flints MM
MM Cos (S)	32.1	6.79	0	trace	0.86	0.81	3200	Synthetic MM with no Fe
MM (Co, Fe) <sub>5</sub> (S-1)	32.3	63.8	φ •	2-10	96.1	0.80	2450	Synthetic MM with 3.9% Fe
* (Co,Fe) <sub>5</sub> (S=2)1 32.3	1 32.3	55.9	11.8	2-10	104.0	0.65	1300	Synthetic MM with 11,8% Fe
MMO.85mo.2 <sup>Co</sup> 5	27.7 MM 6.8 Sm	65.4	1.7	10	98.0	98°0	4950	Modified MM 20% Sm added to natural MM

MM : Synthetic MM.

powder compacts for each composition. Milling for greater than 6 hours increases  ${}_{1}^{H}{}_{C}$  for MMCo $_{4.8}$  but it lowers the B $_{r}$ . Since the six hours milled powder compacts of MMCo $_{4.8}$  alloy show highest value for M $_{r}^{M}{}_{S}$  (0.67) and (BH) $_{max}$  (0.86 MGOe), the powdere compacts of this composition were first tried for sintering studies.

## 7.3.3 Permanent Magnet Properties of Sintered MM-Co Magnets and Their Structure

A systematic sintering study on MM-Co alloy powder compacts was carried out varying the composition, sintering temperature and time. The pulse magnetic field applied during pressing and the compacting pressure were maintained constant at 13 KOe and 14 Tons/cm<sup>2</sup> respectively, for all the pellets used in the sintering studies. The density, Mr/Ms and iHc of the powder compact and its sintered pellets were measured. The structure of the sintered pellets was analysed by X-ray diffraction.

## 7.3.3.1 MMCo4.8 Sintered Pellets

The powder compacts of MMCo<sub>4.8</sub> prepared from 3 h, 6 h and 9 h milled powders were sintered at 1000°C for 15 mins. and their properties are shown in Table 7.4a to 7.4c. The three hours milled powder compact (Table 7.4a) has an iH<sub>c</sub> of 1890 Oe, M<sub>r</sub>/M<sub>s</sub> of 0.52 and a density of 4.9 g/cc. After sintering at 1000°C for 15 mins. the iH<sub>c</sub> decreases to 560 Oe, the M<sub>r</sub>/M<sub>s</sub> drops to 0.13 and the density increases to 6.1 g/cc. Subsequent sintering at higher temperature (1060°C)

TABLE 7.4a PERMANENT MAGNET PROPERTIES OF 3 h MILLED POWDER COMPACTS AND SINTERED PELLETS OF MMCO4.8

Treatment	Mr/Ms	i <sup>H</sup> c (Oe)	Density, g/co
1. Powder compact	0.52	1890	4.9
2. 1000° C/15 min.	0.13	560	6.1
3. 1000° C/15 min. + 1060° C/15 min.	0.09	5 20	7.0
4. 1000°C/15 min. + 1060°C/30 min.	0.08	520	8.0
5. 1000° C/15 min. + 1060° C/ 30 min. + 900° C/1 h	0.06	430	8.0

TABLE 7.4b PERMANENT MAGNET PROPERTIES OF 6 h MILLED POWDER COMPACTS AND SINTERED PELLETS OF MMCO4.8

Treatment	Mr/Ms	i <sup>H</sup> (Oe)	Density,g/cc
1. Powder compact	0.67	2060	4.8
2. 1000° C/15 min.	0.06	520	6.2
3. 1000° C/15 min. + 1060° C/15 min.	0.07	520	7.0
4. 1000° C/15 min. + 1060° C/30 min.	0.06	470	•
5. 1000° C/15 min. + 1060° C/ 30 min. + 900° C/1 h	0.04	4 10	-

TABLE 7.4c PERMANENT MAGNET PROPERTIES OF 9 h MILLED POWDER COMPACTS AND SINTERED PELLETS OF MMCO4.8

Treatment	Mr/Ms	H (Oe)	Density,g/cc
1. Powder compact	0,57	2400	5.0
2. 1000° C/15 min.	0,06	5 20	6.6
3. 1000° C/30 min.	0,05	520	6.7
4. 1000° C/30 min. + 1060° C/15 min.	0,05	5 20	7.0
5. 1000° C/30 min. + 1060° C/ 30 min. + 900° C/1 h	0.03	340	7.6

for longer time (30 m) increases the density to 8.0 g/cc but the M<sub>r</sub>/M<sub>s</sub> and <sub>i</sub>H<sub>c</sub> are further reduced to 0.06 and 430 0e respectively. The same trend is observed for the 6 h and 9 h milled powder compacts. The X-ray pattern of sintered pellet of 9 h milled powder compact is shown in Figure 7.3a. Surprisingly the expected MMCo<sub>5</sub> phase is hardly detectable. The observed lines could be accounted for by 6-Co and MM<sub>2</sub>O<sub>3</sub> phases.

### 7.3.3.2 MMCo4.2 Sintered Magnets

The permanent magnet properties of sintered pellets of MMCo $_{4.2}$  composition are shown in Table 7.5 together with the values obtained for their powder compacts. The sintering was done at 1060°C for 15 m. Both the M $_{r}$ /M $_{s}$  ratio and  $_{i}$ H $_{c}$  decrease on sintering. The X-ray pattern of sintered pellet of 6 h milled powder compact is shown in Figure 7.3b. The

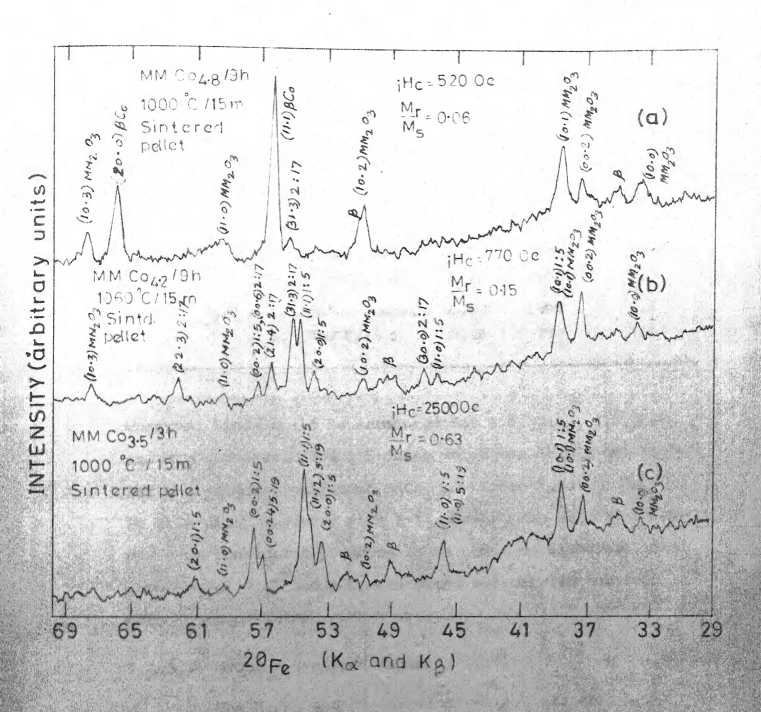


FIG. 7.3 X-RAY DIFFRACTION PATTERNS OF SINTERED PELLETS
OF (a) MMCo4.8 (b) MMCo4.2 AND (c) MMCo3.5 ALLOYS

TABLE 7.5 PERMANENT MAGNET PROPERTIES OF POWDER COMPACTS AND SINTERED PELLETS OF MMCO4.2

Sample	Treatment	Mr/Ms	<sup>Н</sup> с <sup>(Ое)</sup>	Density,g/co
1. MMCo <sub>4.2</sub> /3 h	a. Powder compact b. 1060°C/15 m	0.47	1380 900	gCD.
2. MMCo <sub>4.2</sub> /6 h	a. Powder compact	0.51	1 <b>7</b> 20	4.9
	b. 1060° C/15 m	0.23	950	7.7
3. MMCo <sub>4.2</sub> /9 h	a. Powder compact	0.52	1890	4.8
	b. 1060° C/15 m	0.15	770	7.4

observed lines could be accounted for viz. 2:17, 1:5, 2-Co and MM<sub>2</sub>O<sub>3</sub> phases. The 1:5 phase which was hardly detectable for the sintered pellet of MMCO<sub>4.8</sub> (Figure 7.3a) is present in this alloy together with 2:17 phase. The amount of 2-Co phase is considerably reduced. In order to eliminate the presence of 2:17 and 2-Co sintering was carried out for MMCO<sub>3.5</sub> composition.

## 7.3.3.3 MMCo3.5 Sintered Magnets

The M<sub>r</sub>/M<sub>s</sub> and i<sup>H</sup><sub>c</sub> of sintered pellets of 3 h milled powder compacts of MMCo<sub>3.5</sub> are shown in Table 7.6. Both the M<sub>r</sub>/M<sub>s</sub> and i<sup>H</sup><sub>c</sub> increase for the sintered pellets. The increase in i<sup>H</sup><sub>c</sub> is more than twice the value obtained for its powder compact. These values could be increased further by pulse magnetising the pellets at 60 KOe field.

TABLE 7.6 PERMANENT MAGNET PROPERTIES OF POWDER COMPACTS AND SINTERED PELLETS OF MMCO3.5

Sample	Treatment	Mr/Ms	. <sup>Н</sup> (Ое)	Density,g/co
MMCo <sub>3.5</sub> /3 h	a. Powder compact	0.40	860	- T
	b. 1000° C/15 m magnetised at 11 KOe	0.58	2100	7.6
	c. Pulse magnet- ised at 60 KOe	0.63	2500	7.6
	d. 1000°C/30 m magnetised at 11 KOe	0.58	2100	7.7

The X-ray pattern of this pellet is shown in Figure 7.3c.

The observed lines could be accounted for 1:5, 5:19 and

MM<sub>2</sub>O<sub>3</sub> phases. The 2:17 and R-Co phases are hardly detectable.

Though 1:5 is the predominant phase in the sintered pellet,

still considerable amount of 5:19 is present. To reduce the

amount of 5:19 phase, MMCo<sub>3.8</sub> composition was tried for

sintering.

#### 7.3.3.4 MMCo3.8 Sintered Magnets

The magnet properties for the powder compacts and the sintered pellets of MMCo<sub>3.8</sub> are shown in Table 7.7. Sintering was carried out at three different temperatures 1000°C, 1030°C and 1060°C for 15 m. The 3 h milled powder compact shows M<sub>r</sub>/M<sub>s</sub> of 0.35 and i<sup>H</sup><sub>c</sub> of 1250 Oe. The increase

TABLE 7.7 PERMANENT MAGNET PROPERTIES OF POWDER COMPACTS AND SINTERED PELLETS OF MMCO3.8

Sample	Treatment	M <sub>r</sub> /M <sub>s</sub>	i <sup>H</sup> c (Oe)
MMCo <sub>3.8</sub> /3 h	<ul> <li>a. Powder compact</li> <li>b. 1000° C/15 m, magnetised at 11 KOe</li> <li>c. 1000° C/15 m, magnetised at 60 KOe</li> </ul>	0.35 0.69 0.78	1250 4080 4900
	Powder compact Magnetised at 11 KOe:	0.35	1030
	a. 1000° C/15 m b. 1030° C/15 m c. 1060° C/15 m	0.66 0.61 0.43	2710
	Magnetised at 60 KOe:	0.64	4130
	b. 1030° C/15 m c. 1060° C/15 m	0.70	3400 1800

to 0.78 and 4900 oe respectively for a sintered pellet magnetised at 60 KOe sintering at temperatures higher than 1000°C for 5 h milled powder compacts leads to lower values only. The X-ray diffraction pattern of 1000°C/15 m sintered pellet is shown in Figure 7.4 together with the X-ray pattern of the as-cast MMCo<sub>3.8</sub> alloy. The 5:19, which is the predominant phase in the as-cast alloy, is not detectable in the sintered pellet. Also the MM<sub>2</sub>O<sub>3</sub> peaks not present in the as-cast alloy pattern appears prominently in the X-ray

FIG.74 X-RAY DIFFRACTION PATTERNS OF AS CAST AND
SINTERED PELLETS OF MMCo38 ALLOY

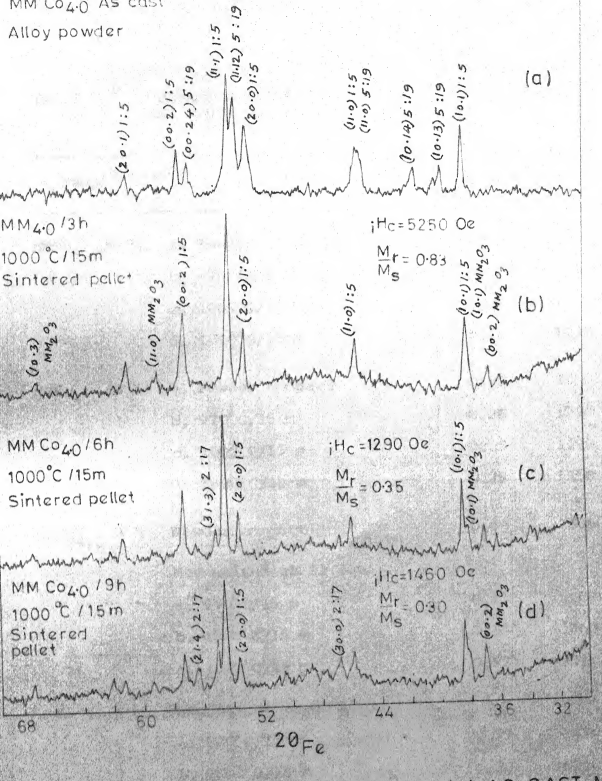
pattern of the sintered pellets suggesting that the oxidation should have occurred during milling or sintering operation.

## 7.3.3.5 MMCo4.0 Sintered Magnets

The MMCo<sub>4.0</sub> alloy was milled for 3 h, 6 h and 9 h and the powder compacts were sintered at 970°C, 1000°C and 1030°C. The M<sub>r</sub>/M<sub>s</sub> and iH<sub>c</sub> values measured are shown in Table 7.8. The 3 h milled powder compact sintered at 1000°C for 15 m shows a maximum M<sub>r</sub>/M<sub>s</sub> of 0.83 and iH<sub>c</sub> of 5250 Oe. The 6 h and 9 h powder compacts show lower values for their sintered pellets. The X-ray patterns of sintered pellets of 3 h, 6 h and 9 h milled powder compacts are compared in Figure 7.5 with the X-ray pattern of as-cast MMCo<sub>4.0</sub> alloy. The as-cast alloy contains 1:5 and 5:19 only whereas the same alloy after milling for 3 hours and sintering at 1000°C for 15 m contains mainly 1:5 and MM<sub>2</sub>O<sub>3</sub>. The sintered pellets of 6 h and 9 h milled powder compacts show some amount of 2:17 phase in addition to the 1:5 and the MM<sub>2</sub>O<sub>3</sub> phases.

7.3.3.6 MM<sub>0.8</sub>Sm<sub>0.2</sub>Co<sub>3.8</sub> and MM<sub>0.8</sub>Sm<sub>0.2</sub>Co<sub>4.0</sub> Sintered Magnets

The MM<sub>O.8</sub>Sm<sub>O.2</sub>Co<sub>3.8</sub> alloy was milled for 3 h and 6 h and sintered at 1000°C for 15 m. On sintering the i<sup>H</sup><sub>C</sub> increased to more than three times their value for powder compacts and the M<sub>r</sub>/M<sub>s</sub> ratio also increased appreciably (Table 7.9). The as sintered pellet of 6 h milled powder compact showed an i<sup>H</sup><sub>C</sub> of 11000 oe. This increased to > 11000 oe on pulse magnetising at 60 KOe. As the peak magnetic field



X-RAY DIFFRACTION PATTERNS OF (a) AS CAST MMCo40 ALLOY AND SINTERED PELLETS OF (b) 3h (c) 6h AND (d) 9h MILLED POWDER COMPACTS OF MM Co40 ALLOY

TABLE 7.8 PERMANENT MAGNET PROPERTIES OF POWDER COMPACTS AND SINTERED PELLETS OF MMCO4.0

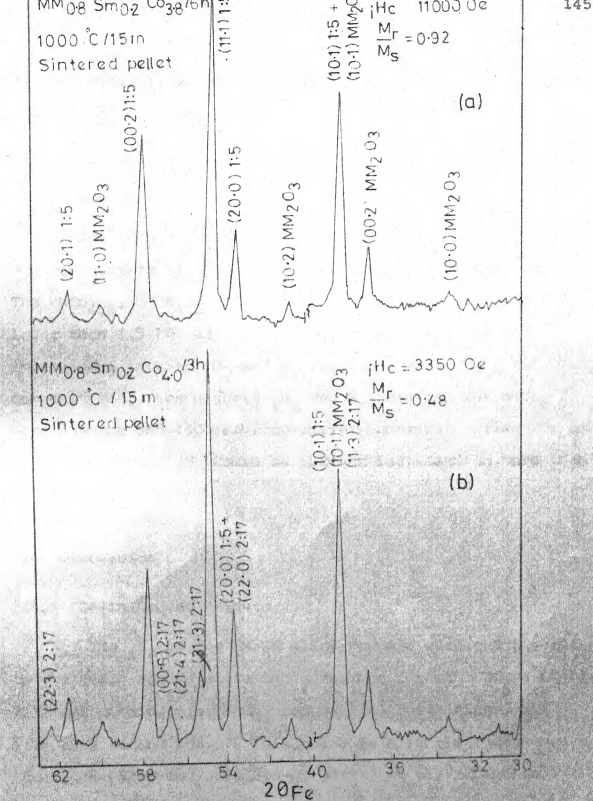
Sample	Treatment	Mr/Ms	i <sup>H</sup> c <sup>(Oe)</sup>
MMCo <sub>4.0</sub> /9 h	a. Powder compact	0.46	2060
	b. 970° C/15 m	0.28	1460
	c. 1000° C/15 m	0.30	1460
	d. 1030° C/15 m	0.26	1290
MMC04.0/6 h	a. Powder compact	0.50	1850
-	b. 970° C/15 m	0.38	1720
	c. 1000° C/15 m	0.35	1290
· ·	d. 1030° C/15 m	0.29	1250
$MMCo_{4.0}/3 h$	Powder compact	0.46	1806
	Magnetised at 11 KOe:		
	a. 970° C/15 m	0.67	3100
	b. 1000° C/15 m	0.67	3960
	c. 1030° C/15 m	0.57	3010
	Magnetised at 60 KOe:		
	a. 970° C/15 m	0.77	3870
	b. 1000° C/15 m	0.83	5250
	c. 1030° C/15 m	0.76	4210

TABLE 7.9 PERMANENT MAGNET PROPERTIES OF POWDER COMPACTS AND SINTERED PELLETS OF MMO.8SmO.2CO3.8 AND MMO.8SmO.2CO4.0

Sample	Treatment	Mr/Ms	i <sup>H</sup> c (0e)
MM <sub>O.8</sub> Sm <sub>O.2</sub> Co <sub>3.8</sub> /3 h	a. Powder compact	0.67	2840
	b. 1100° C/15 m, magnetiged at 11 KOe	0.82	7570
	c. 1000° <b>C/15</b> m, magnetised at 60 KOe	0,90	9720
MM <sub>0.8</sub> Sm <sub>0.2</sub> Co <sub>3.8</sub> /6 h	a. Powder compact	0.66	3270
	b. 1000°C/15 m, magnetised at 11 KOe	0.81	11000
	c. 1000° C/15 m, magnetised at 60 KOe	0.92	>11000
MM <sub>0.8</sub> Sm <sub>0.2</sub> Co <sub>4.0</sub> /3 h	a. Powder compact	0.59	3530
	<pre>b. 1000° C/15 m, magnetised at 11 KOe</pre>	0.48	3354

available for characterising the magnets in the present study was only 11 KOe, the magnitude of its coercivity could not be measured. The  $M_r/M_s$  ratio increased from 0.81 to 0.92 after pulse magnetising. The X-ray pattern shown in Figure 7.6a indicates the presence of 1:5 phase only together with considerable amount of  $MM_2^{0}$ 3.

The  ${\rm MM_{0.8}Sm_{0.2}Co_{4.0}}$  alloy milled for 3 h and sintered at 1000°C for 15 m has lower values for  ${\rm M_r/M_s}$  and  ${\rm i^H_c}$  than its powder compacts (Table 7.9). The study of its



G.7.5 X-RAY DIFFRACTION PATTERNS OF SINTERED PELLETS OF (a) MM<sub>0.8</sub> Sm<sub>0.2</sub> Co<sub>3.8</sub> AND (b) MM<sub>0.8</sub> Sm<sub>0.2</sub> Co<sub>4.0</sub> COMPOSITIONS

X-ray pattern revealed the coexistence of the 2:17 phase with the 1:5 and the oxide phases (Figure 7.6b).

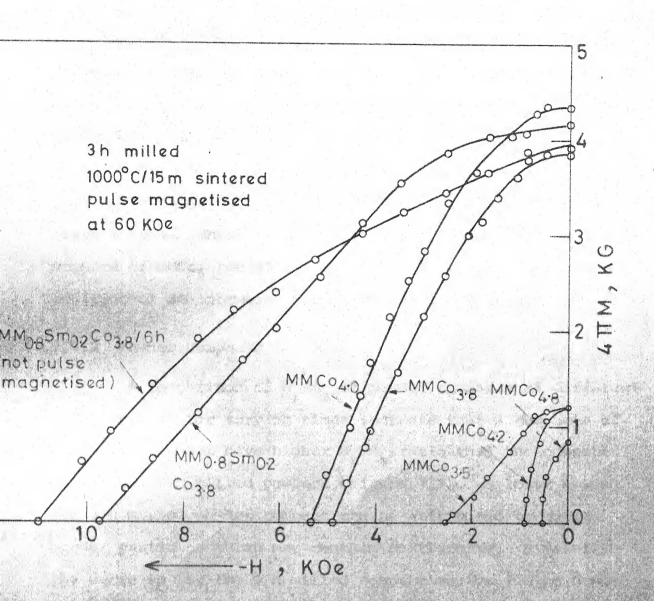
#### 7.3.4 Demagnetisation Plots of MM-Co Sintered Magnets

The intrinsic demagnetisation curves of MM-Co alloys with composition between MMCo $_{3.5}$  and MMCo $_{5.0}$ , all milled for 3 h and sintered at 1000°C for 15 m are shown in Figure 7.7. The MMCo $_{4.8}$ , MMCo $_{4.2}$  and MMCo $_{3.5}$  sintered samples show a Br lower than 1.5 KG and an  $_{i}^{H}{}_{c}$  lower than 3 KOe. A marked improvement in both  $_{i}^{H}{}_{c}$  and Br are seen for MMCo $_{3.8}$  and MMCo $_{4.0}$  compositions. The highest  $_{i}^{H}{}_{c}$  and Br obtained for MMCo $_{4.0}^{H}{}_{c}$  were 5250 Oe and 4400 KG respectively. On substituting 20% Sm to Flints MM the  $_{i}^{H}{}_{c}$  could be further increased to more than 11000 Oe.

#### 7.4 Discussion

#### 7.4.1 Resin Bonded Magnets

The  $_{i}^{H}{_{c}}$  of the MM-Co alloy powders were characterised in the resin bonded state and a maximum  $_{i}^{H}{_{c}}$  of 3200 Oe (Table 7.3) was obtained for MMCo $_{5}$  composition containing 2-10% 5:19 as a second phase. This value is slightly lower than the value (3600 Oe) obtained by Narita et al.  $^{4}$  for MMCo $_{5}$ . The Fe present as impurity in natural mischmetal does not seem to have a deleterious effect on  $_{i}^{H}{_{c}}$  as the MMCo $_{5}$  alloys prepared from the Flints mischmetal and Synthetic mischmetal show same value for  $_{i}^{H}{_{c}}$ . However a small drop in  $_{i}^{H}{_{s}}$  is



G.77 INTRINSIC DEMAGNETISATION PLOTS OF SINTERED MAGNETS OF MMCox ALLOYS 3.5 € X € 4.8

Water was a construction of the

seen for the MMCo $_5$  alloy (Table 7.3) whereas an increase is expected due to the presence of Fe. But MM contains other non-magnetic elements like Mg, Ca etc. upto 1.4 w/o which might decrease the magnetisation. At high concentration of Fe (>2 w/o) the Fe seems to have a deleterious effect on  $i^Hc^*$ . An addition of 20% Sm to MM increases the  $i^Hc$  to a large extent. This was also observed by Nagel  $^2$  for sintered magnets of MMCo $_5$  containing Sm. This increase occurs as the addition of Sm increases the  $^4$ A of the 1:5 phase.  $^2$ 

#### 7.4.2 Powder Compacts

A comparison of  $M_r/M_s$  of powder compacts of different alloys milled for varying times indicate that a compacts of 6 h milled powder have higher M / M ratio than the compact of 3 h and 9 h milled powders (Figure 7.2). A lower value for 3 h milled powder compact can be attributed to coarse powder particles which may contain multigrains. Similarly the decrease for the 9 h milled compact may be due to damage or oxidation of the powder as a result of prolonged milling. The MMCo48 alloy milled for 6 h shows the highest M / M of 0.67. This value is still much less than the expected value (0.8-0.9). The powders in this study were pressed axially in 13 KOe field (of 100-200 millisecond pulse width) in a non-magnetic die where the punch tips were also of non-magnetic material. Application of a continuous magnetic field perpendicular to the pressing direction have been reported to yield good alignment. 7.

#### 7.4.3 Sintered MM-Co Magnets

### 7.4.3.1 The Occurrence of Oxide in MM-Co Sintered Magnets

An oxide of the stoichiometry MM203 could be detected in all the sintered pellets. The MMC04.8 alloy after sintering at 1000°C for 15 m has .2-Co and MM203 with a trace of 2:17. The presence of oxide in sintered Sm-Co8 and MM-Co3 magnets have been reported to an extent upto 0.5 w/o.

Ishigaki et al.8 have made an exclusive study on the oxidation phenomena of SmCo5 during sintering and they report that the iHc of the sintered SmCo5 increases with oxygen content upto 5000 ppm and then decreases at higher concentration. In samples containing 8000 ppm oxygen they could detect Sm2Co17, Sm2O3, CoO and hexagonal Co (co). But in the MM-Co sintered pellets the Co phase existed in f.c.c. form (B) and the oxide had the La2O3 type hexagonal structure. The lattice parameters and the X-ray diffraction data of MM2O3 and 3-Co found in MMCO4.8 sample (Figure 7.3a) are given in Tables 7.10 and 7.11.

#### 7.4.3.2 Compositional Shift in MM-Co Sintered Magnets

The maximum iH<sub>C</sub> obtained for different compositions of sintered MM-Co alloy is plotted in Figure 7.8 as a function of composition. It is seen that the iH<sub>C</sub> reaches a maximum of 5250 Oe for MMCo<sub>4.0</sub> and it falls on either side of this composition. A curve of similar shape has been reported for SmCo<sub>5</sub> sintered magnets. But in that case the maximum iH<sub>C</sub> is observed at SmCo<sub>4.9</sub> (16.9 a/o Sm). Narita<sup>4</sup> in his study

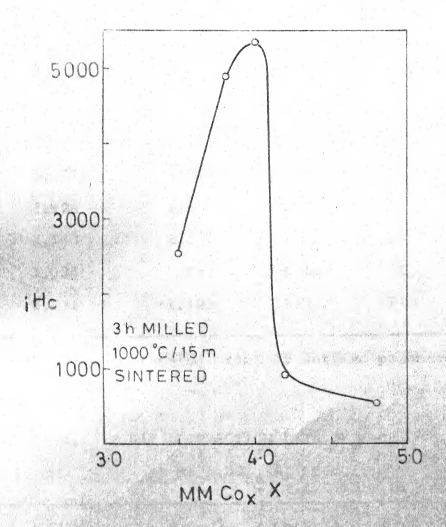


FIG. 7-8 INTRINSIC COERCIVITY VS COMPOSITION FOR SINTERED MMCox ALLOYS 3-8 < x < 4.8

TABLE 7.10 X-RAY DIFFRACTION DATA OF MM $_2$ O $_3$  a: 3.86 $_{71}$  Å; b: 6.05 $_4$ O Å; La $_2$ O $_3$  type hexagonal structure

d <sub>ob</sub> Å	d <sub>cal</sub> Å	(h k,1)	% I <sub>ob</sub>	20 Fe (observed)
3.349*	3.349	1 0.0	28	33.6
3.027*	3,027	0 0.2	31	37.3
2.928	2.930	1 0.1	100	38.6
2.243	2.245	1 0.2	28	51.1
1.936	1.933	1 1.0	31	60.1
1.731	1.728	1 0.3	21	68.1

<sup>\*</sup> Lines taken for calculation of lattice parameters.

TABLE 7.11 X-RAY DIFFRACTION DATA OF 3-Co a: 3.53<sub>56</sub> Å, f.c.c.

d <sub>ob</sub> Å	d <sub>cal</sub> %	(h k 1)	% I <sub>ob</sub>	20 Fe (observed)
2.042	2.041	1 1 1	100	56.6
1.768*	1.768	200	52	66.4
1.250	1.250	2 2 0	32	101.5
1.066	1.066	3 1 1	36	130.5

<sup>\*</sup> Line taken for calculation of lattice parameter.

of sintered MM-Co magnets found a similar shift in composition and reported that MMCo<sub>3.8</sub> after sintering turned out to be single phase MMCo<sub>5</sub> with a maximum iH<sub>C</sub> of 8.3 KOe. The composition of MM used by Narita is very close to the composition of the Indian mischmetal but the former contains 1.9 w/o Sm and 3.9 w/o Fe whereas the Indian mischmetal contains about 6 w/o Fe and no Sm. 10 This shift in phase composition can occur either due to the loss of RE by evaporation during sintering or by selective oxidation of RE. If it is due to evaporation then there should be a weight loss in the sintered pellets and no oxide should be present. But the sintered pellets of MM-Co alloys in this study show negligible weight loss (0.1 w/o) and contain considerable amount of oxide.

The oxidation can occur in one of comminution and sintering operations or in both. In order to determine this the powder compact and its sintered pellets were analysed for their chemical composition. The analysis carried out for  $\text{MMCo}_4$  alloy is shown in Table 7.12.

analysis. (1) The total weight of the elements analysed is always less than 100%. In the case of as-cast alloy the observed deficit of 1% could be attributed to the impurities present in MM that were not analysed but the deficit of 4 to 5 w/o for the powder compacts and the sintered pellets could be accounted for only due to oxidation for which X-ray evidence is there for the sintered pellets. Since the powder

TABLE 7.12 CHEMICAL COMPOSITION OF AS-CAST, POWDER COMPACT AND SINTERED PELLETS OF MMCO4.2

Sample	Composition (w/o)			Total
	Co	RE	Fe	
As-cast	62.6	34.2	2.2	99.0
Powder compact	,			
3 h	<b>5</b> 9.1	34.2	3.0	96.3
6 h	58.0	33.8	3.1	94.9
9 h	58.4	33.6	3.6	95.6
Sintered, 1000° C/15 m				
3 h	59.2	34.4	3.1	96 <b>.7</b>
6 h	58.5	34.1	3.3	95.9
9 h	58.4	33.6	3.6	95.6
. •				

compacts contain particles of few micron size the X-ray peaks were highly diffused and very weak. Hence the presence of oxide could not be confirmed unambiguously in the powder compacts. But the chemical analysis could be used as an indirect proof for it. (2) The compositions of the powder compacts and the sintered pellets are nearly the same suggesting that the factor causing the compositional shift (oxidation) occurs only during comminution and not during sintering. (3) The Fe content of the powder compact is about 1.5 w/o higher than that present in the corresponding as—cast alloy, the amount being higher for the 9 h milled powder than for the 3 h milled powder. This is expected to come

from the wear of the stainless steel rods used as the grinding media. (4) Both for Co and RE the powder compacts show lower value than the as-cast alloy. This is only an apparent decrease and not a real one because the weight of the powder taken for chemical analysis includes the weight of oxygen also which actually goes into water during dissolution in the acid  $(RE_2O_3 + 6HNO_3 - 2RE(NO_3)_3 + 3H_2O)$ . Thus the reference weight taken in the beginning should not be used as such for computing the percent composition of the elements. If we assume a 25-30% oxidation of RE to  $RE_2O_3$  and 5-10% oxidation of Co to CoO and make a mass balance analysis, then this difference in chemical composition between the as-cast and the powder samples could be explained.

#### 7.4.3.3 Structure and Properties of Sintered MM-Co Magnets

Maximum values for the coercivity was observed in those sintered pellets which contained 1:5 phase only. The presence of small amount of 5:19 does not seem to have adverse effect on iHc. But the presence of MM2Co17 which has easy plane anisotropy, needs to be totally avoided to obtain improved iHc. A maximum iHc of 5250 Oe was obtained for the MMCo4.0 composition which contained MMCo5 and MM2O3 only after sintering. Narita obtained a maximum value of 8.3 KOe for MMCo3.8 which after sintering at 1030°C for 30 m contained 1:5 only as determined by thermomagnetic analysis. Owing to the presence of considerable amount of oxide, the 4m Ms of the sintered pellet was 65 emu/g only

whereas a value of 95 emu/g is expected for MMCo $_5$ .  $^{12}$  Consequently the maximum B $_r$  obtained for the oxide containing MMCo $_5$  was 4.5 KG only. Nagel et al.  $^3$  report that a post sintering heat treatment is essential to improve the  $_i^H{}_c$  but in the present study no heat treatment was tried as the sintered pellets contained a large amount of oxide. The minimisation of the oxide content and a suitable post sintering heat treatment should certainly improve the  $_i^H{}_c$  and  $_r^H{}_c$  from the present values.

## 7.4.4 Problem Areas in the Preparation of Sintered MM-Co Magnets

Two main obstacles, oxidation and limitation of magnetic field (of 11 KOe only for characterisation of  $_{i}^{H}_{c}$ ) were found out in this study. The oxidation seems to occur in the pre-sintering stage and mostly during the comminution. The method of powder preparation is to be perfected or to be modified altogether to minimise the oxidation. For MMCo $_{5}$  magnets containing a small amount of Sm, a magnetic field greater than 11 KOe is essential to measure the  $_{i}^{H}_{c}$ .

#### 7.5 Conclusions

A maximum  $_{\rm i}^{\rm H}_{\rm c}$  of 3200 Oe was obtained for MMCo $_{\rm 5}$  powders. On sintering at 1000°C a large scale shift (5 to 6 w/o) in composition was observed as a result of oxidation. MMCo $_{\rm 4.0}$  alloy on sintering at 1000°C for 15 m contained only 1:5

phase together with MM $_2$ O $_3$  and showed an  $_1$ H $_1$ C of 5250 Oe and a B $_1$ C of 4.5 KG. A 20% substitution of Sm for the natural mischmetal increased the value of  $_1$ H $_2$ C to greater than 11 KOe. The oxidation seems to occur during the pre-sintering stage, mostly during the period of powder preparation.

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#### 8. CONCLUSIONS AND SUGGESTIONS FOR FUTURE WORK

Phase relationships in MM-Co-Fe system between 2:7 and 2:17 stoichiometries with Fe content upto 15 a/o were investigated and a 900°C isothermal section of the MM-Co-Fe phase diagram was obtained. The MM-Co-Fe phases between the 2:7 and 1:5 stoichiometries are structurally closely related and show very similar X-ray patterns. Hence extensive thermomagnetic study was carried out in addition to X-ray diffraction and metallography in order to confirm their identities. sequence of occurrence of phases in the binary MM-Co system was found to be essentially similar to that observed in a binary Ce-Co system. MMCo3, MM2Co7, MM5Co19, MMCo5, MM2Co17 and  $\beta$ -Co were found to exist in the MM-Co system between 73 and 92 a/o Co. The 5:19 phase was found to extend upto about 10 a/o Fe into the MM-Co-Fe ternary system whereas the 2:7, 1:5 and 2:17 phases extend to greater than 10 a/o Fe along their respective stoichiometric (MM/Co) lines. A new phase with a T of 340°C was found to coexist with 1:5 and 5:19 phases with a number of alloys between the 5:19 and 1:5 stoichiometric lines. The MM<sub>2</sub>Co<sub>7</sub>, MM<sub>5</sub>Co<sub>19</sub>, MMCo<sub>5</sub> and MM<sub>2</sub>Co<sub>17</sub> were obtained in single-phase form and their standard X-ray diffraction data and crystal lattice parameters were obtained. A homogeneity region for 1:5 phase containing 3 a/o Fe was investigated and it was found to have a narrow homogeneity region at 900°C extending over 1.5 a/o Co. Phase stability

of 1:5 alloy containing different amount of Fe was studied at 700°C in view of their technological importance in the sintering process of these magnets. The MMCo<sub>5</sub> phase was found to undergo an eutectoid type reaction and the increasing amount of Fe substitution (\*\*5 a/o) for Co in the 1:5 phase was found to accelerate the decomposition rate at 700°C.

In order to evaluate the suitability of the MM-Co-Fe phases for permanent magnets, the saturation magnetisation  $(4\pi\,\mathrm{M_S})$  and Curie temperature  $(\mathrm{T_C})$  were characterised for the 2:7, 5:19, 1:5 and 2:17 phases in the MM-Co-Fe system. The MMCo<sub>5</sub> and MM<sub>2</sub>Co<sub>17</sub> phases were found to have attractive properties for both  $4\pi\,\mathrm{M_S}$  and  $\mathrm{T_C}$ . The MMCo<sub>5</sub> phase has a  $4\pi\,\mathrm{M_S}$  of 95 emu/g and the MM<sub>2</sub>Co<sub>17</sub> has a  $4\pi\,\mathrm{M_S}$  of 114 emu/g. Both the 2:7 and 5:19 phases have  $-4\pi\,\mathrm{M_S}$  less than 45 emu/g. The T<sub>C</sub> of the MMCo<sub>3</sub>, MM<sub>2</sub>Co<sub>7</sub>, MM<sub>5</sub>Co<sub>19</sub>, MMCo<sub>5</sub> and MM<sub>2</sub>Co<sub>17</sub> phases were found to be  $-20^{\circ}\mathrm{C}$ , 67°C, 280°C, 540°C and  $-700^{\circ}\mathrm{C}$  respectively. The T<sub>C</sub> of 2:7, 5:19 and 1:5 phases were found to increase with increasing substitution of Fe for Co. From the point of view of  $4\pi\,\mathrm{M_S}$  and T<sub>C</sub> both the MMCo<sub>5</sub> and MM<sub>2</sub>Co<sub>17</sub> seem to be attractive for permanent magnet production.

The addition of Cu to MM-Co alloys was made to know to what extent Cu can be substituted in the MMCo<sub>5</sub> phase with tolerable magnetic properties suitable for permanent magnet production. The study revealed that stable 1:5 phase could be obtained in MM(Co<sub>1-x</sub>Cu<sub>x</sub>)<sub>5</sub> alloys replacing Co with Cu upto 100%. The <u>a</u> and <u>c</u> of the hexagonal 1:5 phase were found

and by during the

to increase linearly with increasing Cu substitution. Both the  $4\pi\,\mathrm{M}_\mathrm{S}$  and  $\mathrm{T}_\mathrm{C}$  of the 1:5 phase were found to decrease steadily with the addition of Cu. It was found that Cu could be substituted upto 15 w/o with a  $\mathrm{T}_\mathrm{C}$  not less than 500°C and a  $4\pi\,\mathrm{M}_\mathrm{S}$  of about 70 emu/g. Beyond 15 w/o Cu, both  $4\pi\,\mathrm{M}_\mathrm{S}$  and  $\mathrm{T}_\mathrm{C}$  were considerably reduced.

MM-Co alloys on the MM rich side of  ${\rm MMCo}_5$  were processed to produce permanent magnets. A selected process parameters: alloy composition, milling time, sintering temperature and sintering time were investigated systematically. The optimum milling time was determined by studying the it of the powder produced rather than its average particle size. A maximum H of 3200 Oe was obtained for the MMCo5 powders. The powder compacts of  ${\rm MMCo}_{\rm x}$  alloys were sintered between 960 and 1030°C. Sintering at temperatures >1000°C was found necessary to obtain high density ( > 90% of the theoretical density) for these alloys. The structure of the sintered pellets was studied closely by metallography and X-ray diffraction. On sintering at 1000°C, a large scale shift in composition (5-6 w/o) was observed. MMCo<sub>4.0</sub> alloy after sintering at 1000°C for 15 minutes contained only 1:5 phase together with MM203. It showed an iH of 5250 Oe and a Br of 4.5 KG. A 20% substitution of Sm in Indian mischmetal increased the 'H to greater than 11 KOe. Chemical analysis and X-ray diffraction study revealed the presence of considerable amount of MM,03 in the sintered pellets. The large scale shift in composition is attributed to oxidation and it seems to occur during the pre-sintering stage, mostly during the period of powder production.

#### APPENDIX 1

SPECIFICATIONS OF RAW MATERIALS: MM, Co, Fe and Cu

Material	Source	Purity/Composition	At. Wt.
Co 1.	Semi Elements, Inc., N.Y., U.S.A.	999%	58,93
2.·	Moldex Co., Bombay, India	99.3%	
Fe	Gallard, Schelsinger Mfg. Co., N.Y., U.S.A.	99.9%	55,85
Cu	Semi Elements, Inc., N.Y., U.S.A.	99.,9%	63.54
ММ	Mischmetal & Flints Pvt. Ltd., Allepy, Kerala, India	Commercial grade Composition: 52.0% Ce, 20.1% La, 15.7% Nd, 4.8% Pr, 6% Fe and 1.4% unidentified impurities	140.82

#### APPENDIX 2

#### COMPUTATIONS OF ALLOY COMPOSITIONS

A procedure followed to prepare an alloy of composition, for example  ${
m RECo}_{4.8}{
m Fe}_{\rm O.2}$  is given.

- Assumption: (a) Total RE in MM = 92.6 w/o
  - (b) Fe in MM = 6 w/o
    - (c) Impurities in MM = 1.4 w/o
    - (d) Effective gm. at. wt.

      of total RE in MM = 140.82 gms.

92.6 gms of RE ≡ 100 gms. of MM

. 140.82 gms of RE  $\equiv \frac{100}{92} \times 140.82 = 152.07$  gms of MM

Fe in 152.07 gms of MM =  $0.06 \times 152.07 = 9.12$  gms

$$REC_{4.8}Fe_{0.2} \equiv MM(C_{04.8}Fe_{x}Fe_{0.2-x})$$

where x = 9.12/55.85 = 0.033

Therefore materials to be weighed:

- (a) 152.07 gms of MM
- (b) 4.8 x 58.93 gms of Co
- (c)  $(0.2 0.033) \times 55.85$  gms of Fe